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(54) Title: EXPANSION AND TRANSDIFFERENTIATION OF HUMAN ACINAR CELLS

(57) Abstract: This invention relates, e.g., to a method for expanding mammalian acinar cells, comprising culturing the cells in a cell culture system comprising a cell culture medium and a cell attachment surface, under conditions wherein the acinar cells undergo a 3-4 fold expansion together with transdifferentiation into a modified cell phenotype (IP cells) showing characteristics of acinar cells and liver cells. The invention also relates to a method for transforming these IP cells to insulin-producing cells *in vitro*, comprising culturing the cells in a novel, defined medium. Also disclosed are suitable culture media for performing these methods, isolated cells having the phenotype of IP cells and/or produced by these methods, and kits for performing the methods.



03/102171

EXPANSION AND TRANSDIFFERENTIATION OF HUMAN ACINAR CELLS

This application claims the benefit of provisional application 60/384,000, filed May 28, 2002, which disclosure is incorporated by reference in its entirety herein.

BACKGROUND OF THE INVENTION

1. Field of the Invention

[0001] The invention relates to compositions and methods whereby, e.g., human pancreatic acinar cells are cultured under conditions that support expansion and transdifferentiation into glandular epithelial cells and subsequently into insulin-producing cells.

2.Background Information

[0002] The potential benefits of taking insulin-producing cells from organ donors and transplanting them into insulin-dependent, Type I diabetic patients is clear. In the Edmonton clinical trials, many patients have lived free from the delivery of exogenous insulin for approximately 2 years after being transplanted with intact islets from organ donor sources. However, current technology requires two organ donor pancreata to generate a sufficient number of islets (about 1 million islets, comprised of about 1,000 cells each) to transplant into one diabetic patient for cellular therapy. Thus, there has been emphasis in the diabetes field to identify new sources of insulin-producing cells for transplantation. Many avenues are being explored, including expansion of islets after harvest and before transplantation and the generation of new islets from stem-like cells derived from the bone marrow, or from precursor cells located in the pancreas. The challenges presented by these approaches are related to maintenance of function of islets over long periods of culture, and of the relative rarity of stem-like cells that can be harnessed for insulin production from the bone marrow and pancreas. The ductular precursor stem-like cells derived from the pancreas are reported to be more efficient than bone-marrow derived cells at differentiation into insulin-producing cells, and this may reflect their site of origin (i.e., pancreas) where they are certainly exposed to many

differentiation signals related to the pancreatic microenvironment. The most abundant cell type in the pancreas is the acinar cell, which comprises about 85% of the pancreas. The acinar cells serve to produce and secrete digestive enzymes and, like islet cells, arise during development from the ductular cell compartment.

[0003] There have been reports that acinar cells, when cultivated *in vitro*, especially under conditions of stress, can undergo a 'transdifferentiation' into a cell type that resembles ductular cells, as determined by expression of CK19, CK7, and carbonic anhydrase (all reputed by the authors to be markers of duct cells) (Kerr-Conte, 1996; WO 02/29010 A2), Hall et al., 1992). Furthermore, Bouwens *et al.* (1998) have shown *in vivo*, in a model of pancreatic duct ligation, that acinar cells in the ligated portion of the pancreas undergo transdifferentiation into cells with a ductular phenotype. Further work has suggested that insulin-producing cells can be produced upon further differentiation of the duct cells in the ligated portion of the pancreas. The acinar cells are also reported to be of limited survivability in primary culture, with some culture conditions leading to loss of at least 50% of cells within a week. While primary duct cells have been demonstrated *in vitro* to convert into insulin-producing cells under some culture conditions (e.g. Bonner Weir, 2000, U.S. Pat. No. 6,011,647), there are no reports of cells that arose from acinar cells *in vitro* differentiating further to produce islet-like cells.

[0004] Prior to the development of the present system, primary pancreatic acinar cells were expanded without differentiation into insulin-producing cells, either in serum-containing medium (undesirable both because of the risk and the uncertainty associated with the use of serum), or in complex serum free media formulations. Likewise, primary pancreatic acinar cells have been transdifferentiated into insulin-producing cells without expansion, producing cells with an insulin-producing phenotype in small numbers. Furthermore, it has not been previously possible to obtain insulin-producing cells in good numbers using acinar cells as starting material.

[0005] Thus, there is a need for a simple cell culture system and method for rapidly generating large numbers of cells that can further differentiate into, e.g., insulin-

producing cells, through expansion and transdifferentiation of the abundant pancreatic acinar cells. Further, there is a need for a cell culture system and method for culturing and transforming such cells into insulin-producing cells. One cell culture system and related method disclosed herein allows a simple, one-step approach that generates expanded cultures that contain at least 80% intermediate progenitor cells that can give rise to insulin-producing cells. A second cell culture system and related method allows the further culturing of these intermediate progenitor cells or other glandular epithelial cells to obtain insulin-producing cells. Both IP cells and insulin-producing cells will be useful for cell-based therapies for the treatment of diseases such as diabetes.

SUMMARY OF THE INVENTION

[0006] The present invention provides compositions and methods whereby, e.g., acinar cells can be cultivated successfully in vitro, undergoing a 3-4 fold increase in cell number over time, and giving rise to a cell population that co-expresses acinar and ductal markers early during the culture (2-3 days ex vivo), then ultimately (e.g., about 7-8 days ex vivo) acquires a modified phenotype characterized by expression of some acinar-associated genes, as well as some liver-associated genes. The genes expressed by these modified cells at about 7-8 days ex vivo include, e.g., ductular cytokeratins (CK7, CK8, CK18 and CK19), hepatic nuclear factor 1 (HNF1), alpha-1 antitrypsin, pi-glutathione s transferase (pi-GST), liver-specific (basic helix-loop-helix (bHLH) transcription factor, Thy-1, CCAAT/enhancer-binding protein (C/EBP)-alpha and C/EBP-beta. These cells exhibit little if any expression of the pancreas-associated genes carbonic anhydrase, cystic fibrosis transmembrane conductance regulator (CFTR), elastase and amylase. By "little if any" expression of a gene is meant herein that gene expression is generally undetectable under conventional methods, such as the hybridization immunocytochemical methods described herein, but expression may be detected by extraordinarily sensitive methods, such as PCR-based analysis. This type of modified cell is referred to herein as an intermediate progenitor ("IP") cell. expanded/transdifferentiated acinar cells (IP cells) can be produced using a general serum-containing media, or, in a preferred method, can be produced without serum on a surface comprising one or more extracellular matrix molecules (ECMs) in the presence of

one or more soluble active factors. ECMs can be presented in 2 dimensional or 3 dimensional culture systems in the presence of soluble active factors.

[0007] The IP cells generated from these cultures are expected to be useful directly in certain medical applications. For example, there is evidence that such cells may under certain conditions become functioning insulin-producing cells when implanted in diabetic patients. The cells can also be used for drug discovery and toxicity studies.

[0008] In addition, according to a further aspect of the invention, the IP cells can be cultivated further, in a serum-free medium composed of any standard serum-free base medium (DMEM:HamsF12, for example) with BSA and combinations of factors, including ECMs, small molecules, and growth factors. After 5-10 days of culture, the IP cells undergo additional steps of differentiation, culminating in the formation of cell aggregates that express pro-insulin and C-peptide. Challenge of these cultures with a high-glucose medium causes release of insulin and C-peptide into the medium, indicating the production in these cultures of functional islet-like cells.

[0009] Thus, in a first aspect, the present invention provides a cell culture system comprising a superior cell attachment surface that also stimulates cellular expansion, and a simple culture medium including effective amounts of one or more soluble active factors, or serum (e.g. fetal bovine serum), added to a base medium composition. The cell culture system will be particularly useful for primary culture of mammalian epithelial cells, particularly human epithelial cells. In a preferred embodiment the cell culture system is used for the expansion and transdifferentiation of primary acinar cells, especially human pancreatic acinar cells.

[00010] The cell attachment surface for this cell culture system is any surface to which the cells can attach and expand, including both 2 dimensional (e.g. plates, flasks, roller bottles, petri dishes, wells etc.) and 3 dimensional (e.g. scaffold) environments. Preferably the surface comprises at least one type of ECM, or a peptide fragment thereof. Cells may, in some circumstances, detach from these surfaces and form self-supporting

aggregates. Suitable fragments include peptides consisting of a sequence of three of more amino acid residues that are identical to any portion of the amino acid sequence of the ECM. Such fragments can be easily made and tested by means known to those of skill in the art. Most preferably the surface is a layer of collagen I. Many other surfaces known in the art are also suitable, such as Collagen VI, Collagen IV, Vitronectin, or Fibronectin. Collagen I is preferred due to ease and cost.

[00011] The base medium to which the soluble active factors are added may be any cell culture medium appropriate for growth and differentiation of epithelial cells. These include, but are not limited to, DMEM, Hams F12, MEM, M-199 and RPMI. The general requirements for such culture media and many suitable examples are known to those of skill in the art. To this base medium is added either serum (such as fetal bovine serum), or a stabilizing protein such as bovine serum albumin (BSA) along with effective amounts of soluble active factors. The medium is preferably serum-free.

[00012] Soluble active factors for the expansion and transdifferentiation of primary pancreatic acinar cells into IP cells include growth factors such as HGF receptor activators and EGF receptor activators. Preferred soluble active factors include one or more of EGF and Transforming Growth Factor-α, IGF1, HGF, betacellulin, prolactin and gastrin 1. HGF, EGF and/or Transforming Growth Factor-α are particularly preferred. Also preferred is the combination of IGF1 and betacellulin.

[00013] In one particularly preferred embodiment, the base medium contains a 1:1 mixture of DMEM and Hams F12. The base medium is completed with the addition of glutamine to a final concentration of ~4 mM, insulin (~0.1-10 μ g/ml, preferably ~0.01 mg/ml), transferrin (~0.5-10 μ g/ml, preferably ~0.0055 mg/ml), selenium (~0.25-5.0 ng/ml, preferably ~0.0067 μ g/ml of sodium selenite), and Epidermal Growth Factor (EGF) (~1-20 ng/ml, preferably ~10 ng/ml); this medium is hereafter referred to as pancreatic cell medium, or PCM. To this base medium formulation, up to ~20% Fetal Bovine Serum (or other serum), preferably between ~10 ~~15% fetal bovine serum, most preferably about 10% or up to about 15% fetal bovine serum) may be added, or, to create

a serum-free culture environment, the following components are added in place of serum: heat-inactivated bovine serum albumin (0.1–2%), Hepatocyte growth factor (HGF) (1-20 ng/ml), and/or Transforming Growth Factor Alpha (TGFa)(1-10 ng/ml). In addition, the medium may contain Betacellulin (0.5-20 ng/ml), Gastrin 1 (0.05-10 ng/ml), Prolactin (1.0–10 ng/ml), and/or IGF-1 (5-100 ng/ml). In particular formulations, greater or lesser amounts of these components may be added in order to achieve a formulation that is effective in supporting the expansion and transdifferentiation of the cells. Persons of skill in the art will appreciate that determining effective amounts of the components will require no more than routine experimentation.

[00014] By the use of this attachment surface and medium, the expansion and transdifferentiation of primary pancreatic cells with the desired phenotype is simplified greatly.

[00015] In a particularly preferred embodiment, the cell culture system is a combination of collagen I coated tissue culture surface (presented in a 2 dimensional or 3 dimensional form) and a serum-free medium containing BSA, insulin, transferrin, selenium, Hepatocyte growth factor (HGF), Epidermal Growth Factor (EGF) and Transforming Growth Factor Alpha (TGFA).

[00016] The cell culture system enables superior attachment in vitro of primary pancreatic epithelial cells for adherent culture compared to prior methods, while creating a cellular environment that promotes expansion of the epithelial component of primary pancreatic cultures with concomitant transdifferentiation of the acinar cells present in the starting material into IP cells, while minimizing emergence of undesired fibroblasts. Advantages of this culture system are ease of construction, few components needed, and that all components are readily available and easily used in the required manner.

[00017] The components of this aspect of the invention may be conveniently packaged in the form of a kit. The kit may include, for example, 1) a cell culture medium such as DMEM: 2) a serum-free medium supplement containing BSA, insulin, transferrin,

selenium, HGF, EGF and TGFA, in suitable amounts to yield the concentrations noted above in the completed medium; and 3) at least one collagen I coated substrate, such as a vessel for tissue culture (e.g., dish(es) with at least one collagen-1 coated tissue culture surface), or collagen-1 coated inserts for use in culture dishes or other laboratory ware. The kit may also optionally include a tissue culture dish or other cell culture accessories and additional reagents that may be required to carry out epithelial cell culture and differentiation.

[00018] Culture systems consisting of scaffolds, collagen coated flasks or other vessels and serum-free base medium may be packaged along with the soluble active factors as a separate vial that would be added to the culture medium just prior to use. The active factor combination can be added to a variety of base media to accomplish the same end, e.g., growth and differentiation of primary pancreatic acinar cells in vitro. Such culture systems should also be useful for other cell types, particularly glandular epithelial cells derived from other organs and tissues, including those from liver, pancreas, intestine, prostate, and breast.

[00019] The collagen I surface provides superior cell attachment (thereby increasing the number of cells that adhere during initial culture and thus enhancing culture efficiency), while the collagen I and the combination of soluble active factors (e.g., HGF, TGFA and EGF) promote continued proliferation of cells over time, leading to an increase in cell number above what has been previously reported for primary pancreatic acinar cells. Furthermore, the expansion of the acinar cells is accompanied by a transdifferentiation in the majority of cells to an IP phenotype, which is potentially a therapeutically useful cell phenotype for the treatment of diseases such as diabetes. This likely occurs due to convergence of the intracellular signaling pathways associated with collagen I, HGF, TGFA and EGF, creating a synergistic response.

[00020] The cell culture system of the present invention has unexpected advantages over systems previously in use. Collagen I, IV, VI, Vitronectin and Fibronectin were expected to enhance cell attachment. However, other extracellular matrix molecules that yielded

equivalent attachment of cells during the initial 18 hours of culture did not promote consistent growth of the cells over time in the serum-free medium containing HGF/EGF/TGFA. The most efficient and cost-effective method of achieving cell expansion AND differentiation into IP cells is to utilize a collagen-I surface and a medium containing reduced serum (preferably less than 20%, more preferably less than 15%, 10%, or 5%, most preferably 2%).

[00021] Another aspect of the invention is a method for culturing mammalian epithelial cells comprising adding said cells to the cell culture system described above, and maintaining them at suitable temperature and atmospheric conditions. By "mammalian epithelial cell" is meant any cell of a tissue or organ with an epithelial cell phenotype, defined by the presence of expression of cytokeratins and often through the presence of markers that suggest a tissue-specific function (i.e., epithelial cells of the skin make keratin, epithelial cells of the intestine make mucin, epithelial cells of the prostate make PSA). In a preferred embodiment, the cells are primary pancreatic cells, particularly human pancreatic cells. Suitable temperature for mammalian cells is usually in the range of about 37°C, but may be varied somewhat according to cell type. The atmosphere can be ordinary air, or other specialized mixtures of gasses suitable for maintaining cells, as will be familiar to persons of skill in the art. Expansion of pancreatic acinar cells can be maximized by decreasing the oxygen tension in the culture atmosphere to less than 21%, while transdifferentiation to IP cells can be enhanced by increasing oxygen tension to greater than 5%. A preferred range of oxygen tension is between about 5% and about 21%.

[00022] In a second aspect, the invention also provides methods and compositions for transforming glandular epithelial cells that have acquired expression of markers characteristic of an intermediate progenitor (IP) phenotype as described above into insulin-producing cells. By "glandular epithelial cell" is meant an epithelial cell that is a component of a gland. Glands are tissues that have a specific function related to secretion of key molecules – most organs in the body have glandular function (liver, intestine, pancreas, prostate, breast, pituitary, adrenal, kidney) whereby they produce and release

hormones, digestive enzymes, or other life-essential fluids. Glandular epithelial cells from endoderm-derived organs (e.g., liver, intestine, pancreas) share many characteristics, including the ability to express many of the same genes. Particularly preferred are glandular epithelial cells from pancreas, for example acinar cells. As used herein, the terms "express" and "expression" generally refer to nucleic acids (e.g., mRNAs) or to protein gene products that are detectable by standard immunocytochemical methods.

[00023] In this aspect, the invention provides a second cell culture system comprising a cell attachment surface and a culture medium that supports and promotes the transformation of glandular epithelial cells into insulin-producing cells. The cell attachment surface is similar to and may be identical to the attachment surface for expanding primary pancreatic acinar cells. It may be presented in the form of a flat surface coated on a vessel or in the form of a scaffold or other surface adapted for cell culture. It can be comprised of, or coated with, any composition that is capable of maintaining cells or supporting cell growth. In a preferred embodiment, it comprises at least one ECM, such as Collagen I, Collagen VI, Collagen IV, Vitronectin or Fibronectin. In a particularly preferred embodiment, the cell attachment surface is Collagen-I.

[00024] In this aspect, the invention provides a further culture medium comprising at least one differentiation promoting factor ("DPF") that promotes the transformation of glandular epithelial cells into insulin producing cells. The DPFs for the transformation of glandular epithelial cells into insulin producing cells can be one or more of Activin A, acidic FGF, basic FGF, C-Natriuretic Peptide (CNP), Calcitonin Gene Related Peptide, Cholera Toxin B Subunit, Dexamethasone, Gastrin-Releasing Peptide, Glucagon-like Peptide-1 (GLP-1), Glucose, IGF1, IGF2, Insulin, Laminin, LIF, Met-Enkephalin, PDGFAA+PDGFBB, Prolactin, Sonic Hedgehog, Substance P, TGF-alpha, Trolox (alpha-tocopherol derivative), or VEGF. Preferred concentrations in culture medium of each of these 23 DPFs are listed in Table 1. Although in some cases one DPF is sufficient, preferably two or more factors are used. As many as all 23 of the factors may be used.

Table 1

Substance	Concentration
	(micrograms/
	mL)
Activin A (human, recombinant)	0.0005
CGRP alpha, (Calcitonin Gene Related Peptide, rat)	0.1905
C natriuretic peptide) (human, porcine, rat: frag 32-	0.10985
53)(CNP)	
Cholera Toxin B Subunit, recombinant	0.0125
DEXamethasone (9 alpha-fluoro-16alpha-	0.002
methylprednisolone)(hydrocortisone analogue)	
FGF acidic (aFGF = FGF1), Recombinant Human	0.0025
GLP-1 (7-36) amide, human (Glucagon-Like Peptide	0.033
1)	
Glucose (base should be l0ow; 0.9 ug/ml)	1.08
Insulin , human (low [] in base media 1 ug/ml)	9.5
LIF, human (leukemia inhibitory factor, human)	0.0025
PDGF AA + PDGF BB MIX	0.005
TGF alpha	0.001
Prolactin (human, recombinant)(a plasma growth	0.0012
hormone)	
Trolox (soluable Vitamin E) (C14H18O4)	0.625
GRP (Gastrin Releasing Peptide, Human)	0.143
IGF-1, recombinant human	0.0025
IGF-2, recombinant human	0.0025
Laminin	2.25
Met-Enkephalin (tyr-gly-glyl-phe-met)	0.003
Sonic Hedgehog (mouse, recombinant)	0.025
Substance P (full length) (H1875 is frag 1-4)	5
FGFb (=FGF2), human Recombinant	0.0025
VEGF	0.0025

[00025] In a preferred embodiment of this aspect of the invention, the culture medium comprises at least one (or as many as all 10) of the following differentiation promoting DPFs:.C-Natriuretic Peptide (CNP), Calcitonin Gene Related Peptide, Cholera Toxin B Subunit, Dexamethasone, Gastrin-Releasing Peptide, Laminin, Met-Enkephalin, PDGFAA+PDGFBB, Sonic Hedgehog, and Substance P.

[00026] In a preferred embodiment, the culture medium that promotes the transformation of glandular epithelial cells into insulin producing cells consists of a 1:1 mixture of DMEM and Hams F12 plus the components listed in Table 2. This medium is sometimes referred to herein as "Media or Medium G9."

Table 2

	Factor	Substance	Concentration (Final) ug/ml	
1	ANP	Atrial Natriuretic	0.1530	
		Peptide, Rat (28 amino acids)	0.1050	
2	CCK-8-frag-amide	CCK8, Fragment 26-33 Amide (Cholecystokinin) (Asp-Tyr(SO3H)-Met-	0.0250	
		Gly-Trp-Met-Asp-Phe- NH2)		
	Caerulin sulfate	Caerulein (Pyr-Gin- Asp-Tyr(SO3H)-Thr- Gly-Trp-Met-Asp-Phe- NH2	0.0300	
4	Cholera Toxin-B	Cholera Toxin B Subunit, recombinant	0.0125	
5	Dex	DEXamethasone (9 alpha-fluoro-16alpha- methylprednisolone)(h ydrocortisone analogue)	0.0020	
6	FGF-7	FGF7 (KGF)	0.0025	
	GLP-1	GLP-1 (7-36) amide, human (Glucagon-Like Peptide 1)	0.0330	
8	GRP	GRP (Gastrin Releasing Peptide, Human)	0.1430	
	Gastrin-1,	Gastrin I Human	0.0000	
	Glucose	Glucose (base should be l0ow; 0.9 ug/ml)	1.0800	
	HGF	Hepatocyte Growth Factor (HGF) recombinant	0.0025	
12	IGF-1	IGF-1, recombinant human	0.0025	
	IGF-2	IGF-2, recombinant human	0.0025	
	Insulin	Insulin (low [] in base media 1 ug/ml)	9.5000	
	Leu-Enkephalin	Leu-Enkepthalin (tyr- gly-gly-phe-leu)	0.0030	
16	Nicotinamide	Nicotinamide	610.0000	

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17	PTHRP-frag-1-34	pT II RP (Parathyroid	0.2060
- 1		Hormone Related	1
		Peptide (1-34), human)	
18	Progesterone	Progesterone	0.0030
19	Prolactin	Prolactin (human,	0.0012
		recombinant)(a plasma	
1		growth hormone)	
20	Retinol acetate	Retinoic Acid (Vitamin	0.0250
		A)	
21	SHH	Sonic Hedgehog	0.0250
		(mouse, recombinant)	
22	Sodium selenite	Selenium (Selenous	0.0250
		Acid, Na salt)	
23	Soybean trypsin Inhibitor	Trypsin Inhibitor,	0.5000
	· ·	soybean (type I-S)	
24	TGF-beta sRII	TGF beta sRII (soluable	0.0050
		receptor type 2)	
25	Transferrin	transferrin	2.7500
26	Trolox	Trolox (soluable	0.6250
		Vitamin E) (C14H18O4)	
27	VIP	Vasoactive Intestinal	0.0665
		Peptide (VIP), human	
28	aFGF	FGF acidic (aFGF =	0.0025
		FGF1), Recombinant	
		Human	
29	bFGF	FGFb (=FGF2), human	0.0025
		Recombinant	
30	n-Butyric acid	n Butyric Acid, Sodium	4.5400
		Salt	
		•	

[00027] The components of this aspect of the invention may also be conveniently packaged in the form of a kit. The kit may include, for example, 1) a cell culture medium such as DMEM, Hams F12, or a combination thereof; 2) a serum-free medium supplement containing: BSA and the DPFs Activin A, acidic FGF, basic FGF, C-Natriuretic Peptide (CNP), Calcitonin Gene Related Peptide, Cholera Toxin B Subunit, Dexamethasone, Gastrin-Releasing Peptide, Glucagon-like Peptide-1 (GLP-1), Glucose, IGF1, IGF2, Insulin, Laminin, LIF, Met-Enkephalin, PDGFAA+PDGFBB, Prolactin, Sonic Hedgehog, Substance P, TGF-alpha, Trolox (alpha-tocopherol derivative), or VEGF, or two or more of these components in combination, in suitable amounts to yield the concentrations noted in Table 1 in the completed medium; and 3) tissue culture dish(es) with at least one collagen-1 coated tissue culture surface (or collagen-1 coated inserts for use in culture dishes or other laboratory ware). The kit may also optionally include a tissue culture dish and/or other cell culture accessories and additional reagents that may be required to carry out epithelial cell culture and differentiation. In other

embodiments, the kit may contain any of the media or media components discussed herein.

[00028] Culture systems consisting of scaffolds, collagen coated flasks or other vessels and serum-free base medium may be packaged along with the DPF(s) as a separate vial that would be added to the culture medium just prior to use. The DPF combination can be added to a variety of base media to accomplish the same end, e.g., growth and differentiation of primary pancreatic acinar cells in vitro. Such culture systems may also be useful for other cell types, particularly other epithelial cells derived from glandular tissues, including those from liver, pancreas, intestine, prostate, and breast.

[00029] The invention also provides a method for converting glandular epithelial cells into insulin-producing cells comprising culturing the glandular epithelial cells in the cell culture system described above. The method may further comprise removing the culture medium from the cell culture, re-feeding the cell culture with a serum-free medium with glucose, and measuring proinsulin production C-peptide production, or insulin release.

[00030] Furthermore, the invention provides an isolated population of insulin-producing cells containing cytoplasmic granules with immunodetectable proinsulin, insulin, and/or c-peptide that is derived from a population of cells of which a subset of cells expressed at least one marker associated with IP cells (e.g., expressed some acinar-associated genes, as well as some liver-associated genes, including, e.g., ductular cytokeratins (CK7, CK8, CK18 and CK19), HNF1, alpha-1 antitrypsin, pi-glutathione s transferase (pi-GST), liver-specific bHLH transcription factor, Thy-1, C/EBP-alpha and C/EBP-beta, and expressed little if any of the pancreas-associated genes carbonic anhydrase, cystic fibrosis transmembrane conductance regulator (CFTR), elastase and amylase).

[00031] By an "isolated" cell or population of cells is meant herein that the cell or cell population is removed from its original environment (e.g., the natural environment if it is naturally occurring), and isolated or separated from at least one other component with which it is naturally associated. For example, a naturally-occurring cell present in its natural living

host is not isolated, but the same cell, separated from some or all of the coexisting materials in the natural system, is isolated. Such cell or cell populations could be part of a cell culture or cell population, and still be isolated in that such culture or population is not part of its natural environment.

[00032] In one preferred embodiment, the insulin-producing cells are derived from glandular epithelial cells obtained from mammalian pancreas, such as primary acinar cells.

[00033] The data disclosed in the examples below are generated from freshly isolated human pancreatic cells. The expansion of primary human pancreatic cells in these conditions produces cultures with a mixed epithelial IP phenotype, suitable for *in vitro* studies of IP cells for a variety of purposes, and suitable for transplantation *in vivo* for cell therapy for the treatment of diseases such as diabetes. The IP cells generated by these methods may also be useful in the study of pancreatic cell biology, as normal controls in the study of pancreatic epithelial cancers, and to test the effects of drugs/compounds on normal pancreatic epithelial cells (ductal or acinar). Furthermore, the cells may be further cultured to yield insulin-producing cells as demonstrated below.

BRIEF DESCRIPTION OF THE DRAWINGS

[00034] Figures 1A-D show microscopic images after treatment of starting material with antibodies to amylase (Fig. 1A), insulin (Fig. 1B), and CK19 (Fig. 1C) and the composition of the cell pellet of freshly isolated primary human pancreatic cells (Fig. 1D).

[00035] Figure 2 shows growth curves constructed from primary human pancreatic cultures grown in commercial medium (with serum) or in the described pancreatic cell medium (PCM) with serum.

[00036] Figure 3 shows a comparison of cell expansion in the base medium composition described vs. base medium + soluble growth factors (serum-free formula) vs. base medium + fetal bovine serum.

[00037] Figures 4A-B shows the effect of different culture surfaces on total cell number (Fig. 4A) and cell phenotype (Fig. 4B) after expansion.

[00038] Figures 5A-B show a comparison of cell phenotype after expansion in serum-containing (5A) and serum-free (5B) medium containing all soluble active factors.

[00039] Figure 6 shows high power images of cell cultures expanded in various conditions, including serum-free base media supplemented with 3 soluble active factors, HGF, EGF & TGFA. Note epithelial morphology.

[00040] Figure 7 shows a demonstration of growth of IP cells on ECM-coated surfaces as determined by metabolic activity assay over time. Note superior growth when Collagen I surface is combined with the media formulation described herein, yielding results superior to the combination of Matrigel and commercial media with serum.

[00041] Figure 8A (upper left) shows expression of amylase by acinar cells after two days of culture (red staining), Figure 8B (lower left) shows expression of CK19 (green staining), Figure 8C (right) shows an overlay of the two images, showing co-expression (yellow) in a large proportion of cells.

[00042] Figure 9 shows changing phenotype of primary acinar cells in culture over 5 days. Amylase is red, CK19 is green. Note appearance of yellow (amylase + CK19) on Day 2 and 3.

[00043] Figures 10A and 10B show primary human pancreatic cells that were expanded in serum-containing medium on Collagen I coated surface. Images were analyzed to

determine total cells (Figure 10A, blue nuclei) and total positive cells (Figure 10B, blue nuclei surrounded by green staining for CK19).

[00044] Figure 11 shows light microscopic (200X) appearance of pancreatic acinar cells cultured on a collagen I surface with all DPFs (Activin A, 0.5 ng/ml; acidic FGF, 2.5 ng/ml; basic FGF, C-Natriuretic Peptide (CNP), 0.11 μg/ml; Calcitonin Gene Related Peptide, 0.19 μg/ml; Cholera Toxin B Subunit, 12.5 ng/ml; Dexamethasone, 0.002 μg/ml; Gastrin-Releasing Peptide, 0.143 μg/ml; Glucagon-like Peptide-1 (GLP-1), 0.033 μg/ml; Glucose, 1.08 μg/ml; IGF1, 0.0025 μg/ml; IGF2, 0.0025 μg/ml; Insulin, 9.5 μg/ml; Laminin, 2.25 μg/ml; LIF, 0.0025 μg/ml; Met-Enkephalin, 0.0030 μg/ml; PDGFAA+PDGFBB (0.0050 μg/ml: 0.0025 μg/ml of PDGFAA + 0.0025 μg/ml PDGFBB); Prolactin, 0.0012 μg/ml; Sonic Hedgehog, 0.025 μg/ml; Substance P, 5.0 μg/ml; TGF-alpha, 0.0010 μg/ml; Trolox (alpha-tocopherol derivative), 0.625 μg/ml; and VEGF, 0.0025 μg/ml).

[00045] Figure 12A (top right panel) shows immunocytochemical analysis with CK19 antibodies (green).

[00046] Figure 12B (lower right panel) shows immunocytochemical analysis with C-peptide antibodies (red).

[00047] Figure 12C (left panel) shows an overlay image demonstrating the colocalization of CK19 and C-peptide (orange). Blue portions are DAPI stained nuclei.

[00048] Figure 13A shows insulin release upon glucose challenge in IP cells that have not been detached and relocated (subcultured) during the growth and differentiation process.

[00049] Figure 13B shows insulin release upon glucose challenge in IP cells that have been subcultured according to Example 10.

[00050] Figure 13C shows C-peptide release upon glucose challenge in IP cells that

have not been subcultured according to Example 10.

[00051] Figure 14 shows the Insulin/DNA ratio in subcultured and nonsubcultured cells

that are treated with Combinations 1, 2 and 3 of DFP media, as described in Example 11.

[00052] Figure 15 shows insulin release in response to base level glucose (5 mm) and a

glucose challenge (22mm) over 10 days of culture in PCM and DPF media, as described

in Example 13.

[00053] Figure 15A shows insulin release in response to base level glucose (5 mm) and

a glucose challenge (22mm) over 14 days of culture in PCM and DMG9 media, as

detailed in Example 14.

[00054] Figure 16 is a graphical representation of the characteristics of the 17 classes of

genes shown in Table 6, as indicated in the last column of the Table, as detailed in

Example 14.

DETAILED DESCRIPTION OF THE INVENTION

[00055] In describing preferred embodiments of the present invention, specific

terminology is employed for the sake of clarity. However, the invention is not intended

to be limited to the specific terminology so selected. It is to be understood that each

specific element includes all technical equivalents, which operate in a similar manner to

accomplish a similar purpose. Each reference cited here is incorporated by reference as if

each were individually incorporated by reference.

[00056] The following abbreviations are used:

BSA: bovine serum albumin

BMP Bone Morphogenetic Protein

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bHLH: basic helix loop helix

DMEM: Dulbecco's Modified Eagle's Medium

TGF β 1: Transforming Growth Factor β 1

ECM: extracellular matrix molecules; naturally occurring proteins produced by cells of a tissue that provide structural support as well as a source of cellular signals related to adhesion. Examples are collagen, vitronectin, fibronectin, laminin.

EGF: Epidermal Growth Factor

Hams F12: Ham's Nutrient Mixture F12

HGF: Hepatocyte growth factor

HNF-1: Hepatic nuclear factor 1

IGF1: Insulin-like growth factor 1

IGF-II: Insulin-like growth factor 2

IP cells: Intermediate progenitor cells derived from an epithelial cell, such as, e.g., a pancreatic acinar cell or a liver cell, wherein the derived cells express some acinar-associated genes, as well as some liver-associated genes, including, e.g., cytokeratins (CK7, CK8, CK18 and CK19), HNF1, alpha-1 antitrypsin, pi-glutathione s transferase (pi-GST), liver-specific bHLH transcription factor, Thy-1, C/EBP-alpha and C/EBP-beta, and express little if any of the pancreas-associated genes carbonic anhydrase, cystic fibrosis transmembrane conductance regulator (CFTR), elastase and amylase).

PDGF-A: Platelet derived growth factor alpha

PDGF-B: Platelet derived growth factor beta

TGFA, TGF- α : Transforming Growth Factor α

[00057] As used herein, the term "culture system" is intended to mean a system for growing and/or differentiating cells in culture, which comprises a cell attachment surface, preferably one that also stimulates cellular expansion, and a culture medium, which includes effective amounts of one or more factors, or serum (e.g. fetal bovine serum), added to a base medium composition.

[00058] When referring to active soluble factors and DPFs herein, "effective amount" means an amount that either alone or in combination with other included factors is

effective in promoting either expansion and differentiation into IP cells, or into insulinproducing cells, as applicable.

EXAMPLES

I. Expansion and transdifferentiation of primary acinar cells into glandular epithelial cells (Culture Phase I)

Materials and methods:

[00059] Starting Material: Primary human pancreatic acinar cells are collected as waste from standard COBE gradient preparation of islet cells for transplantation (Lake et al., 1989). After density gradient centrifugation, the islets are present as a layer between 1.063 density and 1.10 density, and the remaining cells are collected as the pellet that sediments to the bottom of the gradient based on density. Approximately 48 hours after collection of the cells at the transplant center are received by the inventors in non-tissue-culture treated polystyrene flasks and are suspended in RPMI + 10% fetal calf serum at a density of approximately 2.0 million cells/ml. Cell number and viability is assessed by trypan blue exclusion and enumeration on a hemacytometer by light microscopic observation.

[00060] Phenotypic Evaluation of Starting Material. A preparation of starting material was formalin fixed and paraffin-embedded as a cell pellet about 24 hours after initial harvest of the pancreatic cells. Paraffin sections were prepared, placed on slides, and subjected to immunocytochemical analysis with antibodies to insulin (Biogenex, San Ramon, CA), CK19 (Biogenex), and Amylase (Biogenex). A minimum of (3) sections per sample were assessed with each marker. All antibody staining was carried out according to the manufacturer's suggestion with pre-diluted commercial antibodies. For CK19, a 3 minute treatment with pepsin enzyme (Biogenex) preceded the blocking step for the purpose of antigen retrieval. Briefly, the sections were rehydrated through graded ethanols, followed by a 15 minute incubation in phosphate buffered saline (PBS) without calcium and magnesium. Protein Blocker (Biogenex) was added for 30 minutes prior to addition of primary antibodies. After (3) 5-minute washes, biotinylated secondary

antibody (Biogenex) was added at a 1:100 dilution and sections were incubated for 30 minutes at room temperature. After (3) 5-minute washes, Alexa488 or Alexa-596-conjugated StreptAvidin (Molecular Probes, Eugene, Oregon) was added for fluorescent visualization. For each slide, a minimum of (3) 200x images were captured using a Nikon fluorescent microscope fitted with a SPOT camera (Diagnostic Systems, Inc., Webster, TX). The images were assessed quantitatively using image analysis software (MetaMorph/Universal Imaging Corporation, Downington, PA) to determine relative fraction of insulin-positive, CK19+, and amylase+ cells. Insulin+ cells are the beta cells of the islets, CK19+ cells are the primary ductal cells, and amylase+ cells are the acinar cells (see Example 1).

Example 1: Characterization of cell culture conditions

A. Serum-free medium

[00061] Freshly isolated primary human pancreatic cells were collected as a pellet from a COBE cell separator, fixed in formalin, paraffin-embedded, sectioned, and analyzed with antibodies to amylase, CK19, and Insulin. Images (Figures 1A and 1B) were collected on a Universal Imaging System (Universal Imaging Corporation) and analyzed with MetaMorph Software. This cell pellet (Figure 1C) was comprised of 1.0% insulin+cells (beta cells of the islet), 5.8% CK19+ cells (primary ductal cells), and 93.2% amylase+ and unlabeled (acinar cells and other cell types).

[00062] Primary human pancreatic cells were then seeded at 10⁴ or 10⁵ cells/ cm² onto tissue culture treated polystyrene in either DMEM commercial medium plus 10% fetal bovine serum or in PCM plus 10% fetal bovine serum. Replicate cultures were harvested at 3 day intervals via trypsinization and live cells (as determined by trypan blue exclusion) and enumerated on a hemocytometer. The results (shown in Figure 2) demonstrate that the (serum-containing) medium formulation described herein is superior to commercial media formulation for growth and maintenance of primary pancreatic cells. Figure 3 compares the results of expanding the cells for 6 days in base medium, base medium plus all of the soluble active factors [HGF, ~1~20 ng/ml, preferably ~5.0 ng/ml; TGFA, ~1~10 ng/ ml, preferably ~2 ng/ml; Betacellulin, ~0.5~20 ng/ml,

preferably ~10ng/ml; Gastrin 1, ~0.05~10 ng/ml, preferably ~0.06 ng/ml; Prolactin, ~1.0~10 ng/ml, preferably ~2.4 ng/ml; and IGF1, ~5~100 ng/ml, preferably ~ 5 ng/ml] and base medium plus 10% serum. The serum-free media formulation meets/exceeds expansion provided by media + serum.

[00063] The cell expansion experiment was repeated essentially as above, except that the base medium was supplemented with only three of the soluble active factors: TGF, HGF, and EGF. Figure 6D compares the results of expanding the cells in the various media; Figures 6A, 6B and 6C show high power images of the cell cultures expanded under the various media conditions.

B. ECM Surfaces

[00064] The attachment of primary human pancreatic cells was evaluated by counting the number of attached cells vs. the number of cells initially seeded on a panel of ECM surfaces comprised of Collagen I (1 µg/cm²), Fibronectin (3 µg/cm²), Laminin (2 μg/cm²), Vitronectin (1 μg/cm²), Matrigel (1 μg/cm²), Human ECM (1 μg/cm²), or Poly-D-Lysine (3 µg/cm²). In one condition, a mixture of Collagen IV, Laminin, and Fibronectin was utilized. ECMs were placed into solution at the above concentrations and allowed to coat tissue culture-treated polystyrene surfaces according to manufacturer's suggestions of 1 hour at room temp. Excess ECM solution was then removed and surfaces were rinsed twice in water. Just before seeding cells, the water was aspirated, then cells were seeded onto the ECM surface at a density of 1 x 105 cells/cm2 in growth medium (PCM) composed of DMEM:HamsF12 mixture (1:1) with 4mM glutamine, 1x ITS supplement (GIBCO 51500-056), 10% Fetal Calf Serum (Inactivated, Qualified, GIBC 26140-079), and 10 ng/ml Epidermal Growth Factor (EGF) (BD 4001). Cells were seeded onto tissue-culture polystyrene surface as a control. After 18 hours, unattached cells were washed away and remaining attached cells were re-fed with PCM and allowed to grow for 7 days prior to evaluation. Cultures were fixed in 10% formalin and subjected to immunocytochemistry with antibodies for CK19 and Amylase as described previously to determine phenotypic composition. Cells were counterstained with DAPI fluorescent blue nuclear stain to visualize individual cell nuclei for cell counting. The metabolic

activity of cells subjected to the various conditions was determined by an MTS assay. Viable cells were measured using the MTS assay (Promega CellTiter 96 Aqueous One Solution Cell Proliferation Assay), a colorimetric method for determining the number of viable cells in proliferation or cytotoxicity. The results of this analysis are shown in Figure 7.

Example 2: Further studies with ECM surfaces and various media components

[00065] Primary pancreatic cells, composed of >90% non-islet/non-duct cells, were plated onto various coated surfaces at a density of 28,900 cells/well (10⁵ cells/cm²). Unattached cells were washed off after 18 hours, and cultures were re-fed and allowed to grow for 8 days. Cultures were fixed in formalin (10%) and subjected to phenotypic analysis with antibodies to CK19 and Amylase. The results are shown in Figure 4A-B. While Collagens I, IV, Laminin, Fibronectin, and Matrigel provide a suitable surface for cell attachment and expansion, maintenance of acinar (amylase+) phenotype along with the presence of an increased proportion of cells with a glandular epithelial phenotype (CK19+) was superior on Collagen I. More than 50% of cells analyzed expressed amylase and more than 50% of cells analyzed expressed CK19, suggesting that a subpopulation of cells in these experimental conditions express both markers.

[00066] Tissue culture-treated polystyrene culture surfaces were coated with Collagen I as described above. Tissue culture medium (PCM) was prepared as described above. In some cases, serum was replaced with Fraction V BSA (99% pure, heat inactivated, Sigma), along with combinations of soluble growth factors, including IGF1, IGF2, betacellulin, HGF, EGF, and TGF-alpha. Optimal seeding density is between 10⁴ and 10⁵ cells/cm², as demonstrated in Example 3. Cells were seeded onto collagen-coated flasks (150 cm²) at 1.5 x 10⁶ cells/flask in PCM. After an ~18hr attachment period, unattached cells were washed away with gentle aspiration/rinse, followed by re-feeding with fresh medium. Cultures were monitored over time by metabolic assay (MTT) and by trypsinization and cell counting, to establish cell number (see example 3). Cell phenotype at the end of the culture period was assessed as follows: small-scale cultures were set up simultaneously in 96-well plates. At the end of the culture phase, monolayer cells were

fixed in 10% formalin for a minimum of 1 hour. After formalin was removed and monolayers were rinsed, cultures were subjected to immunocytochemistry as described in previous section for CK19, amylase, insulin, and vimentin (a marker of fibroblasts). The relative fraction of CK19+ cells was determined by quantitative image analysis as described above (see Example 4). After formalin was removed and monolayers were rinsed, cultures were subjected to immunocytochemistry as described in previous section for CK19 and vimentin (a marker of fibroblasts). Cells were also stained with amylase antibodies, but did not produce positive results due to release of digestive enzymes, such as amylase, by the cells over time in culture. The relative fraction of CK19+ cells was determined by quantitative image analysis as described above (see example 4). Acquisition of ductal markers by acinar cells was verified by demonstrating concomitant expression of CK19 and amylase in cell subpopulations during days 2-3 of culture (see example 5). For these experiments, CK19 primary antibodies were reacted with formalinfixed cell cultures, followed by visualization with Alexa488-conjugated Goat anti-mouse IgG (Molecular Probes). Then, cells were subjected to a blocking step (Protein Blocker, BioGenex), followed by application of the second primary antibody (anti-amylase). Visualization of the amylase was accomplished by application of Alexa594-conjugated Goat Anti-Mouse IgG. Images were collected as described above. At the end of a 7-day culture period in the conditions described herein, between 65-90% of the cells in the culture express CK19, while less than 20% express vimentin (see example 6). Variations in the relative proportion of CK19+ cells probably reflect heterogeneity due to age, gender, and other unique characteristics of individual patients.

Example 3: Density of cell seeding

[00067] Primary pancreatic cells were seeded at (3) densities on tissue-culture treated polystyrene dishes (60mm) and fed with PCM. Light microcopic observations were made daily. At the 24-hour timepoint, dishes were sacrificed and stained with trypan blue to assess viability. The results are shown in Table 3.

TABLE 3

Seeding Density	At 24 Hours:	At 48 Hours:	After 3 days
			Growth:
10 ⁴ cells/cm ²	Most cells attached,	Mitotic Figures present	Epithelial
	trypan blue negative	(light microscopy)	monolayer
	(live)		forming
10 ⁵ cells/cm ²	Most cells attached,	Mitotic Figures present	Epithelial
	trypan blue negative	(light microscopy)	monolayer
	(live)		forming
10 ⁶ cells/cm ²	Some cells attached,	Few Mitotic Figures	Cells are
	most are trypan blue positive	present (light microscopy)	detached; some
	(dead)		fibroblasts present

[00068] Example 4. Cells were grown on a Collagen I surface, at 37°C in 21% oxygen, in PCM medium or in base medium with 2 % BSA, 2 ng/ml TGF-α, 10 ng/ml EGF, and 10 ng/ml HGF. After 7 days, cultures were fixed in 10% formalin and subjected to immunocytochemical analysis with fluorescent detection, followed by automated image collection and analysis. The results are shown in Figures 5A and 5B. Fibroblast (vimentin+) fraction, glandular epithelial cell fraction (CK19+), and fraction of unlabeled cells (Other) are similar after expansion. This suggests that replacement of serum with the serum-free medium maintains fraction of CK19+ cells without overgrowth of fibroblasts as compared to cells grown in serum-containing media.

[00069] Example 5. Primary pancreatic acinar cells were cultured for several days in a 1:1 ratio of DMEM and HamsF12, with 10% fetal bovine serum, 0.01 mg/ml insulin, 0.0055mg/ml transferrin, 0.0067µg/ml sodium selenite, 10 ng/ml EGF, 4mmol/liter glutamine and antibiotics. After 2 days of culture (4 days ex vivo), expression of amylase by the acinar cells is still strong (Figure 8A, upper left panel, red staining) as determined by immunocytochemistry. Expression of CK19 is also apparent (Figure 8B, lower left panel, green staining). Overlay of the two images (Figure 8C) demonstrates clear co-expression of amylase and CK19 in a large proportion of the cells, indicating that an

intermediate cell exists from active conversion of amylase+ acinar cells to an amylase+/CK19+ mixed acinar/ductal phenotype (AD cells). Daily evaluation of cultures (Figure 9) demonstrated that onset of CK19 expression begins around Day 2 of culture and by Day 5 cultures have lost most immunodetectable amylase expression and CK19 expression is prevalent.

[00070] Example 6. After 7 days of growth in PCM / Collagen I surface, cells were fixed, stained with antibodies to CK19, and counterstained with nuclear DAPI. Total cell number was evaluated by automated image analysis (Figure 10A left panel, blue-stained cell nuclei), while CK19+ cells were counted (Figure 10B, right panel, green-stained cell cytoplasm). Of 378 total cells, 342 were immunopositive for CK19 (90%). After approximately 7 days of culture using conditions described herein, the acinar cells have concrete ductular characteristics, now referred to as IP cells. For most primary human cultures, more than 80% of cells in the culture after about 7 days express markers such as CK19 that are associated with ductular cells from a variety of tissues.

[00071] Example 7: Gene Expression Analysis of 7-Day Cultures (IP Cells).

Two independent IP cell cultures were subjected to Clontech 8K Atlas Gene Array analysis. IP cells were obtained by culturing primary acinar cells in a cell culture system comprising PCM and a Collagen I surface. Monolayer cultures were rinsed 2x with PBS, then detached from the flasks with 0.25% trypsin. Cells were pelleted by centrifugation at 1,200 RPM for 3 minutes in a swinging bucket centrifuge. Cell pellets were resuspended and washed 2x in PBS before a final centrifugation at 1,200 RPM for 3 minutes as described above. The supernatant was discarded and gently aspirated to remove as much liquid as possible from the cell pellet, which was then quick-frozen in a dry-ice/ethanol bath and stored at -80°C until transfer to BD Clontech where gene expression analysis was performed, using conventional techniques.

[00072] Labelled P-33 cDNA probes were prepared from the 30 μ g of total RNA from each sample by first enriching for poly A + RNA using a streptavidin-magnetic bead separation method that is part of the Atlas Pure Total RNA Labeling system. The labeled

probes from each sample were hybridized with the plastic human 8 K gene arrays for about 16 hours, the arrays were washed and imaged according to the Atlas array protocols. The Atlas image 2.7 software was used to align array images with the array grid template and to exclude false background signals or false signals due to strong signal bleedover. The transcript signals were then extracted from these aligned arrays using the Atlas Image 2.7 software and further statistical analysis of the changes in gene expression were performed.

[00073] In general, mRNA transcription was assayed, by hybridization to suitable oligonucleotide probes. In a few cases, e.g., for CK19 and amylase, the protein expression product was measured, using conventional methods of immunohistochemistry. A summary of the expression by these cell populations of a selected set of genes is presented in Table 4. Table 4 contains a list of genes expressed in IP cells and a comparison of expression patterns in primary acinar cells and primary ductal cells. Gene products identified as "+" were expressed; those identified as "++" were strongly expressed. Gene products designated ® are found in regenerating pancreas.

TABLE 4:

Gene	IP Cells	Primary Ductal Cells	Primary Acinar Cells
Aquaporin 1	+	++(mRNA)/+(protein)	0
Aquaporin 5	++	+(mRNA)/-(protein)	
Aquaporin 8	+	-	+
Insulin Receptor Substrate-2	+	++ ®	
Protein Kinase B (AKT)	0	+	+
Calpain (mu)	++		:++
CFTR (Cystic Fibrosis	0	0	0
Transmembrane			
Conductance Regulator)			
Claudin 27	nd .	++	- /
Claudin 3	+	+	+
Claudin 4	+	+	+
Claudin 5	nd	. • . • . • . • . • . • . • . • . • . •	++
Carbonic Anhydrase II	0	0	0
Inositol 1,4,5 triphosphate	++	-	++
Rc, Type 3		•	
MUC-1	+	++	+

MUC-6	++	+	
Cytokeratin 7	++	++	•
Cytokeratin 8	++	+	
Cytokeratin 18	+	+	
Cytokeratin 19	+	++	-
alpha v integrin	o	+	
alpha 3 integrin	++	n/a	
beta 3 integrin	0	+	
beta 4 integrin	+	n/a	
beta 5 integrin	0	+	
fibronectin	0	+	
collagen IV	+	+	
vitronectin	0	+ ®	
MMP2	0	Trace	
MMP9	+	Trace	
TIMP1	++	Trace	
TGF-alpha	Trace	++ ®	
Gastrin	0	++ ®	
ICAM-1	0	0	
ICAM-2	0	0	
ICAM-3	0	0	
Pancreatitis Associated	++	+ ®	Trace
Protein (PAP)		ai e	
Reg-1	+	0	Trace
pyrimidinergic receptor P2Y	+	+	
Syndecan	+	+(low)	Trace
Glutathione S Transferase -	+ 0	+	7,
pi		4	

II. Transformation of glandular epithelial cells into insulin producing cells — Generating insulin-producing cells by differentiation of IP cultures (Culture Phase II)

[00074] IP cultures can be utilized to generate insulin-producing cells by placing the cells in a second phase of culture that includes a surface, such as Collagen I, that promotes attachment of the IP cells combined with a defined medium formula that lacks serum but contains combinations of the following differentiation promoting factors: Activin A, acidic FGF, basic FGF, C-Natriuretic Peptide (CNP), Calcitonin Gene Related Peptide, Cholera Toxin B Subunit, Dexamethasone, Gastrin-Releasing Peptide,

Glucagon-like Peptide-1 (GLP-1), Glucose, IGF1, IGF2, Insulin, Laminin, LIF, Met-Enkephalin, PDGFAA+PDGFBB, Prolactin, Sonic Hedgehog, Substance P, TGF-alpha, Trolox (alpha-tocopherol derivative), and VEGF. In the following examples, the base medium is composed of a 1:1 mixture of HamsF12 and DMEM with antibiotics and 0.2% Bovine Serum Albumin (Fraction V, heat inactivated 99% pure). In one example (Combination 1), the base medium contained Cholera Toxin B, Dexamethasone, GRP, GLP-1, Glucose, IGF-1, IGF-2, Insulin, Prolactin, Sonic Hedgehog, Trolox, aFGF, and bFGF. In another example (Combination 2), the base medium contained Activin A, CGRP-alpha, CNP, Glucose, GLP-1, IGF-2, Insulin, LIF, Met-Enkephalin, Prolactin, Sonic Hedgehog, aFGF, and vEGF. In a third example (Combination 3), the base medium contains Activin A, CGRP-alpha, Cholera Toxin B, Dexamethasone, Glucose, GLP-1, Insulin, LIF, Laminin, Met-Enkephalin, PDGFAA/BB, Sonic Hedgehog, Substance P, TGF-alpha, aFGF, and VEGF. The concentrations of these media supplements are listed in Table 1.

[00075] AD cells were placed into culture by either: 1) trypsinizing the cells from the surface on which they were generated, and redistribution onto a fresh attachment-promoting surface at a density of ~5 x 10⁴ cells/cm², or 2) removing the medium, washing 2x in PBS to remove traces of old medium, and cultures re-fed with the new medium (described above) containing differentiation promoting factors. Cells are cultured for a period of 4-10 days at 37°C and 21% oxygen. On Day 5, half of the medium is removed and replaced with an equal volume of fresh medium containing differentiation promoting factors.

Phenotypic Analysis of IP cells After Differentiation Culture.

[00076] Morphological assessment of IP cells cultured in differentiation conditions described above was captured by light microscopy (see Example 8, below). The cellular phenotype of the cells comprising these cultures was assessed by immunocytochemistry as described above using monoclonal antibodies to vimentin, pro-insulin, C-peptide, MUC-1, and CK19 (See Example 10, below). Briefly, cultures were fixed with 10%

formalin for 1 hour at room temperature, then washed with PBS and subjected to immunocytochemical protocol. (See Example 9, below).

Functional Analysis of IP cells After Differentiation Culture.

[00077] The ability of the aggregated cell clusters to release insulin and C-peptide was assessed by subjecting the cultured cells to a glucose challenge as follows. Cells that had been cultured in differentiation medium for 7-10 days were washed 3x in PBS, then refed with either 1) base medium (described above) with 5mM Glucose, or 2) base medium with 22mM glucose. After 18 hours, the cell-conditioned medium was collected and subjected to ELISA analysis for insulin and C-peptide release (Diagnostic Systems Laboratories (DSL)). ELISAs were conducted using the standard range assay procedure according to manufacturer's specifications. Plates were incubated on a shaker during the assay and results were read in a Tecan spectrophotometric plate reader. Total ng of insulin or C-peptide per well were calculated for each media condition, for both 5mM glucose media and 22mM glucose media (See Example 10).

[00078] Example 8. Pancreatic acinar cells were cultured in Base Medium + ITS + Serum (10%) for 1 week, then trypsinized (treated with 0.25% Trypsin without EDTA for 10 minutes at 37°C) and transferred to a fresh collagen-1 coated surface and placed in a medium containing all 23 DFPs listed. Over a period of 3-5 days, the cells readily formed three-dimensional pod-like structures, clearly observable by light microscopy (Figure 11). Some larger pods detached from the culture surface after about 4-6 days in culture, and remained viable, as determined by trypan blue exclusion. The pod-like structures were hypothesized to be aggregations of insulin-producing cells, and subjected to further analysis as described below.

[00079] Example 9. Pod-like structures, generated the same manner as described in the previous example, were fixed in 10% formalin and subjected to immunocytochemical analysis first with CK19 monoclonal antibodies, then with C-peptide monoclonal antibodies, as described above. Figure 12A shows a group of cells (DAPI stained nuclei are blue), some of which are immunopositive for CK19 (green staining). Figure 12B

shows the same group of cells, many of which are positive for C-Peptide, which is produced when the proinsulin molecule synthesized within the cell is cleaved to yield mature insulin; the C-peptide stained cells are red, with a typical granular staining of the cytoplasm. Figure 12C shows a higher power overlay image, demonstrating colocalization of CK19 and C-peptide in a small subset of cells. Co-stained cells appear yellow-orange on the overlay image.

[00080] Example 10. Cells cultured in base medium (negative control), or in Combinations 1, 2 and 3 of the differentiation promoting media, were evaluated for their ability to release insulin and C-peptide into the culture medium. In addition, we assessed whether increasing concentrations of glucose led to the release of a greater quantity of insulin and C-peptide, indicating an islet-like functionality. First, the cells were cultured for 1 week in base medium + EGF(10 ng/ml) + ITS + 10% fetal bovine serum (PCM). Then, cells were either subjected to a wash and medium change (non-subcultured), or to a wash, trypsinization/detachment, reseeding, and medium change. Replicate cultures were re-fed with either base medium (serum-free), fresh PCM, or one of the three combinations of differentiation promoting media (all serum-free). After 10 days, differentiation media were removed, cultures were washed 3x with PBS, then re-fed with serum-free base medium containing either 5mM glucose or 22mM glucose (final concentration). After 18 hours, the conditioned media were collected and subjected to ELISA analysis with antibodies to either Insulin or C-Peptide (DSL laboratories). Figures 13A, 13B and 13C, respectively, show insulin release by nonsubcultured cells, and insulin release and C-peptide release in response to glucose challenge. Since some of the cultures contain insulin, and cells can take insulin up from the medium, production of Cpeptide is an important confirmation that the cells are synthesizing insulin de novo from the synthesis and processing of proinsulin. Furthermore, the production of insulin and Cpeptide is increased in most cases with increasing glucose concentration, suggesting an islet-like function of cells within these cultures. Note that little insulin or C-peptide is produced in the base medium that contains no DPFs.

[00081] Example 11. Both the quantity of insulin and the quantity of DNA were measured in IP cells subjected to differentiation culture with or without enzymatic detachment and subculturing. Cultures were carried out precisely as described in the previous paragraph. DNA was measured utilizing a standard Picogreen assay (Molecular Probes), while insulin was measured by ELISA assay. Total ng of Insulin was divided by total μ g of DNA in the sample, thus providing the insulin:DNA ratio value, in order to calculate a ratio of the quantity of insulin present vs. the number of cells present (reflected by DNA content). The results are shown in Figure 14. In each of the differentiation media combinations, the insulin:DNA ratio is increased compared to base medium, suggesting that more insulin is produced on a per cell basis in the presence of DPFs than when cultured without them. Furthermore, the insulin:DNA ratio is increased slightly in some conditions upon glucose challenge (22mM glucose vs. 5mM), suggesting that the cells respond to glucose by releasing a greater quantity of insulin.

[00082] Example 12. Insulin-producing cells obtained by the preceding method were subjected to gene expression analysis as described above. Table 5 contains a list of the highest expressed genes, their position on the Clontech atlas 8K gene array, and relative expression of these genes (after normalization). Table 5 is attached hereto as Appendix 1

[00083] Example 13. Primary human pancreatic cells were seeded at 0.5 x 10⁵ cells/cm² in PCM on a collagen-1 surface and grown for 7 days. Insulin was measured at Days 1, 7, and 10 as follows: Growth medium was removed, wells were washed 3x in phosphate buffered saline. After a pre-incubation for 1 hour at 37C in base medium without insulin, with 5mM glucose, media was removed and replaced with either 1) base medium (without insulin) with 5mM glucose, or 2) base medium (without insulin) with 22mM glucose. Insulin was measured in cell-conditioned media after 18 hours at 37° C. After 7 days of culture, PCM medium was replaced with either 1) fresh PCM, 2) serumfree base medium, 3) serum free base medium with all 23 differentiating factors, 4) serum-free combination 1, or 5) serum-free combination 2. The results are shown in Figure 15. After 3 days exposure to the differentiating factors, increased insulin release is noted in presence of differentiating factors. The results on Day 1 argue against the

presence of a significant number of insulin-producing cells in the starting material, demonstrating the *de novo* generation of insulin-producing cells from acinar cells in the primary culture. It can be seen in the Figure that at the end of 10 days, insulin release in response to a glucose challenge is much greater in the DFP media than in the PCM or base medium, verifying the stimulatory effect that the DFPs exert on transformation of glandular epithelial cells into insulin-producing cells.

[00084] Example 14. Human pancreatic acinar cells were cultured on a collagen I surface in PCM from Day 1 to Day 7, thus generating a culture of IP cells at Day 7. On Day 7, the IP cells were washed and the PCM medium was replaced with the G09 differentiation medium containing the 30 factors listed in Table 2. At each time point (Days 1, 7, 10 and 14), insulin release was measured by washing the cultures three times with PBS, then challenging the cultures with a 1:1 mixture of DMEM and HAMs F12 containing either 5mM or 22mM glucose. After 18 hours of exposure to the glucose, supernatants were collected and insulin measured by ELISA. The results are shown in Figure 15a.

III. Expression studies at several time points of primary human acinar cells that are expanded, allowed to differentiate into IP cells and then allowed to differentiate further into insulin-producing cells

[00085] Example 15. Three independent samples of primary human pancreatic acinar cells were seeded and expanded described above. From Day 0 to Day 8, cells were on collagen I surface, seeded at 10⁴ cells/cm², in PCM. On Day 8, the medium was changed from PCM to the medium with the active factors shown in Table 2. Cells were fed twice with G09 (50% of medium replaced) between days 8 and 16. The cells remained on the surface throughout the culture process. Cultures were harvested at 3 days after the initial plating (actively trans-differentiating acinar cells), 8 days after plating (IP cells) and 16 days after plating (putative insulin producing cells) and subjected to gene expression analysis, as described in Example 7. mRNA expression data were obtained with 12K microarrays from Clonetech.

[00086] Briefly, growth medium was removed from the culture flasks and cells were lysed in trizol LS (Invitrogen) chaotrope/phenol reagent for about 2 minutes by pipetting the lysis solution over the cell layer. Three ml of RNAse free water was added per 9 ml of lysis solution in an Oak Ridge Cetrifuge tube. 2.4 ml chloroform was then added and the solution vigorously vortexed for 1 minute. The aqueous and organic phases were then separated by cetrifugation at 4°C and the upper aqueous phase containing RNA was removed to a clean PET tube. The RNA was precipitated by isopropanol precipitation, washed with 70% ethanol and redissolved in 200 µl of RNAse free water. A chaotrope lysis reagent was immediately added to the RNA and it was further purified using a Qiagen spin column method with a DNAse digestion step. The purified RNA was finally eluted in 80µl RNAse free water and stored at -80°C.

[00087] Labelled P-33 cDNA probes were prepared from the 30 μ g of total RNA from each sample by first enriching for poly A + RNA using a streptavidin-magnetic bead separation method that is part of the Atlas Pure Total RNA Labelling system. The labeled probes from each sample were hybridized with the plastic human 12 K gene arrays for about 16 hours, the arrays were washed and imaged according to the Atlas array protocols. The Atlas image 2.7 software was used to alighn array images with the array grid template and to exclude false background signals or false signals due to strong signal bleedover. The transcript signals were then extracted from these aligned arrays using the Atlas Image 2.7 software and further statistical analysis of the changes in gene expression were performed.

[00088] The raw expression data were analyzed as follows: (1) We filtered out genes that were not expressed at any of the 3 conditions/time points; (2) We normalized all of the microarrays against each other to remove differences from array-to-array and the effects of variability in sample processing, hybridization, etc.; (3) We identified genes which showed a statistically significant difference among the conditions/time points; and (4) We clustered the genes based on their temporal patterns in a way that is consistent with the design of the study and the changes in phenotype.

[00089] Table 6 shows expression data for the genes that were identified by the above analysis. This Table is attached hereto as Appendix 2. These identified genes were expressed at high levels at both Day 3 and Day 8, or their expression increased substantially between Day 3 and Day 8. The Table also shows the expression levels of these genes at Day 16, and the mean expression for all three condition/time points. The Table also shows the ratios of expression at various times: "I to A" is the ratio of expression of putative insulin-producing cells (Day 16) to acinar (Day 8) cells; "Int to A" is the ratio of IP cells (Day 8) cells to acinar cells (Day 3).

[00090] The data shown in Table 6 were further analyzed by clustering them into one of 17 "classes," whose features are summarized on the Table. A graphical representation of the characteristics of these 17 classes in presented in Figure 16.

[00091] The data from the Day 8 time points in Table 6 were also grouped with regard to whether the genes expressed at Day 8 in these cells belong to the classes of genes expressed normally in (1) liver and pancreas; (2) pancreas-associated genes; (3) liver-associated genes; or (4) progenitor-associated genes. The results are shown in Table 7.

Table 7	BDT Intern	BDT Intermediate Cells	s		
Genes Expressed in Liver and Pancreas	Sample 1	Sample 2	Sample 3	Hepatic Pattern of Expression	Pancreatic Pattern of Expression
CK18	+	+	+	hepatic lineage	acinar cells
CK8	+	+	+	hepatic lineage	acinar cells
CK19	+	+	+	bile duct	duct cells
CK7	+	+	+	bile duct	duct cells
HNF1	+	+	+	liver tc factor	in beta cells
α -1 antitrypsin	‡	‡	‡	produced in differentiated hepatocytes	yes
Notch-1		,	Trace	Involved in differentiation of cells into hepatic lineage	developing pancreatic epithelium
α-fetoprotein		•		developing hepatocytes and hepatic progenitors	developing pancreatic ducts
Notch-3	Trace	Trace	Trace	involved in differentiation of cells into hepatic lineage	pancreatic mesenchyme and endothelium
Notch-4	Trace	Trace	Trace-	involved in differentiation of cells into hepatic lineage	pancreatic mesenchyme and endothelium
Jagged-2	+	+	+	involved in differentiation of cells into hepatic lineage	yes
pi-glutathione s transferase (pi-GST)	+	+	‡	developing liver	duct cells and centroacinar cells
y-glutamyl transferase				developing bile duct	acinar cells
Pancreas-Associated Genes	Sample 1	Sample 2	Sample 3	Hepatic Pattern of Expression	Pancreatic Pattern of Expression
carbonic anhydrase	ı		Trace	ou	duct cells
CF transmemb conductance regulator	,		Trace	OL.	duct cells
elastase	•		•	OU	acinar cells
amylase	•	1	1	OU	acinar cells
insulin	+	Trace	Trace	OU	islet
somatostatin	Trace	Trace	Trace	OU	islet
Pancreatic Polypeptide	+	+	Trace	OU	islet
Glucagon		•	ı	OL	Islet

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Liver-Associated Genes	Sample 1	Sample 2		Sample 3 Hepatic Pattern of Expression	Pancreatic Pattern of Expression
Sialyltransferase-6				produced in differentiated hepatocytes	no
Liver-specific bHLH transcrip factor	+	+	+	liver-specific Tc factor	no
Thy-1	+	+	+	hepatic oval cell marker	no
Glucose-6-phosphatase	,	. •	Trace	hepatic lineage, progenitors and adult	No
Glutamine synthetase	1	1	Trace	hepatocyte	No
Carbamoyl phosphate synthetase-1		,	,	hepatocyte	no
Dipeptidylpeptidase IV	•	•	,	hepatocyte	OU
C/EBP-a	‡	‡	+	liver-specific Tc factor	*turned on and upregul during hepatization of pancreas (copper-deficient diet)
C/EBP-beta	‡	‡	‡	liver-specific Tc factor	*turned on and upregul during hepatization of pancreas (copper-deficient diet)
Progenitor Cell-Associated Genes	Sample 1	Sample 2	Sample 3	Tissue	
Musashi-1	ı	,	٠	Intestine	
Nestin		•	,	Pancreas / Neuronal	
CD34	•	,	•	Hematopoletic	
Thy-1	•	1	,	Hepatic progenitors	
BMP-2	•		,	Neuronal	
BMPRcIA	•	•	Trace	Mesenchyme	
o-kit	•		,	Liver / Pancreas / Neuronal	
chromogranin A	Trace	Trace	Trace	Neuroendocrine / Liver / Intestine	
PDX-1	ı	•	+	Pancreas	

[00092] As can be seen, at Day 8 IP cells no longer expressed genes consistent with pancreatic acinar cells, nor did they express a complement of genes specific for pancreatic ductular cells. The IP cells expressed low levels of some markers associated with pancreatic islets, including insulin, somatostatin and pancreatic polypeptide, suggesting that at least some cells in the population are competent to express endocrine genes of the pancreatic islets.

[00093] Surprisingly, the IP cells also expressed several liver-specific transcription factors (e.g., C/EBP alpha, C-EBP-beta) and other markers of mature and developing liver, including low levels of Thy-1, a marker associated with hepatic "oval" stem cells. This suggests that the differentiating cells were not moving simply from pancreatic acinar to pancreatic ductal, but had developed into a cell with both hepatic and pancreatic characteristics, while not fitting into any single gene expression profile of one of these cell types. The cells generated in this example resemble the cells that emerge from the pancreas of rodents that are fed a copper-deficient diet. (See, e.g. Rao et al., 1988). The pancreas of such animals goes through an acute phase of pancreatitis followed by "hepatization" of liver (which means cells that begin to express hepatic genes rather than pancreatic genes). Liver-like cells have also been reported in human fetal pancreas (Tsanadis et al., 1995) Isolated cells generated by the methods of the present invention (e.g., by propagating primary acinar cells or other types of endodermal cells or progenitor cells by the methods of the invention) are to be distinguished from naturally occurring cells that may have some of the characteristics of IP cells, such as oval cells or cells isolated from the pancreas of a rodent on a copper-deficient diet.

[00094] Cells having the characteristics of these IP cells may be useful for, e.g., therapeutic approaches in the treatment of diabetes. Furthermore, although the cells in this example were derived from pancreas, other epithelial tissues, or perhaps even any endoderm-derived tissue, may provide additional sources of cells that can be differentiated into cells having a similar phenotype. Suitable tissue types include, e.g., liver or intestine. These IP cells express genes associated with pancreas, liver, intestine and neuronal tissues. For example, they express mucin, CK19 and CK7, which are

common markers associated with duct cells in the pancreas, liver and intestine. Thus, the gene expression pattern seen in these IP cells may serve as a predictive measure for cells derived from each of these tissues for the purpose of generating insulin-producing cells. Furthermore, IP cells may, under appropriate conditions, give rise, not only to pancreatic islet cells, but also to hepatocytes or any endoderm-derived tissue.

[00095] The disclosures of the following references, cited above in part, relate to the present invention:

WO 02/29010 A2 (Kerr-Conte);

Bonner-Weir, S. et al., Proc. Natl. Acad. Sci. USA 97: 7999-8004 (2000),

Bouwens, L., Microsc. Res. Tech. 43: 332-6 (1998),

Bowens, L. et al. Diabetologia 41:629-33 (1998);

Gmyr, V. et al., Diabetes 49:1671-80 (2000);

Gmyr, V. et al. Cell Transplant 10:109-21 (2001),

Gmyr, V. et al. Diabetes 49:1671-80 (2000),

Hall, P.A. et al., J. Pathol. 166: 97-103 (1992);

Kerr-Conte, J. et al., Diabetes 45:1108-14 (1996);

Kerr-Conte, J. et al., Transplant Proc 27:3268 (1985);

Pattou F. et al., Bull. Acad. Natl. Med. 184:1887-99 (2000);

Rao, MS et al Biochem Biophys Res Comm. 156:131-6 (1988);

Rooman, Ilse et al., Diabetes 51: 686-90 (2002);

Rooman, I, et al. Diabetologia 43:907-14 (2000);

Rooman, I. et al., Gastroenterology 121: 940-9 (2001);

Trivedi, N. et al. Endocrinology 142:2115-22 (2001);

Tsanadis, G. et al. Histol. Histopathol. 10:1-10 (1995);

Wang, R.N. et al., Diabetologia 38:1405-11(1995);

USP 6,011,647 (Ammon Peck).

[00096] The embodiments illustrated and discussed in the present specification are intended only to teach those skilled in the art the best way known to the inventors to make and use the invention, and should not be considered as limiting the scope of the

present invention. The exemplified embodiments of the invention may be modified or varied, and elements added or omitted, without departing from the invention, as appreciated by those skilled in the art in light of the above teachings. It is therefore to be understood that, within the scope of the claims and their equivalents, the invention may be practiced otherwise than as specifically described.

[00097] The entire disclosure of all applications, patents and publications, cited above and in the figures are hereby incorporated in their entirety by reference.

GENE aquaporin 5 actin, beta actin, beta	Position on Atlas 8K Array Re 7547 3952 8176 8346	Relative Expression, Sample 1 4539 3598 3047	Relative Expression, Sample 2 4550 3442 3747
promin i special AT-rich sequence binding protein 1 (binds to nuclear matrix/scaffold- associating DNA's)	3035	1437	986
cardiac-specific homeo box CCAAT/enhancer binding protein (C/EBP),	2050 1588	1395 1374	2850 1671
Deca RAS guanyl releasing protein 2 (calcium and DAG-regulated)	8262	1263	1616
cartilage paired-class homeoprotein 1 paired-like homeodomain transcription factor 1	6805 6805	1166 1113	1241 756
CTITY OF BOOK NOW	7621	1063	801
complex)	1000 i	† 1	2 6
	5613	935	808
ATPase, Ca++ transporting, plasma membrane 2	0467	9 9	8/8
ficolin (collagen/fibrinogen domain- containing) 3 (Hakata antigen)	. 824	901	829
nositol polyphosphate phosphatase-like 1	2989	832	425
protein tyroslne phosphatase, receptor type, S	8367	806	810
integrin, alpha 3 (antigen CD49C, alpha 3 subunit of VLA-3 receptor)	1743	758	647
	6851	743	658
-	7135	725	, 852
	7436	402	. 1617
midkine (neurite growth-promoting factor 2)	2470	682	805
	3147	678	3084

antityopsin), member 1, and procein asset, and procein asset, retined 6 protein coupled receptor antityosein, member 1, and procein asset, retined 6 protein coupled receptor (seizure- response factor 1 (EGF-response factor 1) 7325 630 646 646 6460 factor 1) 7325 630 646 646 6460 factor 1) 7325 733 733 733 733 733 733 734 6401 factor protein) 7325 733 733 733 733 733 733 733 733 733 73	calbindin 2, (29kD, calretinin) serine (or cysteine) proteinase inhibitor,	7000 5214	676 668	786 678
Se 7325 630 7583 589 7583 589 7583 589 7069 558 6323 558 6489 558 7903 500 6872 428 4202 4428 4202 4411 1965 6604 405 6604 405 713 370 7491 368	ciade A (aipna-1 anuproteinase, antitrypsin), member 1 extra G anticl counted recentor	707	853	969
58	reurial of protein coupled receptor myosin regulatory light chain 2, smooth muscle isoform	2499	638	561
7583 589 7583 589 7583 589 3593 576 7069 558 6323 558 4576 536 4093 516 6489 503 7903 500 6872 428 4202 441 1965 407 6604 405 3577 395 370 2720 370 7491 368	butyrate response factor 1 (EGF-response factor 1)	7325	630	646
7583 589 3593 576 7069 558 6323 558 4576 536 4093 516 6489 503 7903 500 6872 428 4202 411 1965 407 6604 405 3577 395 370 2720 370 7491 368	ie receptor (seizure-	7583	589	394
3593 576 7069 558 6323 558 4576 536 4093 516 6489 503 7903 6872 428 4202 411 1965 407 6604 405 2439 391 713 375 7491 368	ne receptor (seizure-	7583	589	394
6323 558 6323 558 4576 536 4093 516 6489 503 7903 500 6872 428 - 4692 428 - 411 1965 407 6604 405 6604 405 2439 391 713 375 2720 370 7491 368	peptidase enhancer	3593	576	713
6323 558 4576 536 4093 516 6489 503 7903 500 6872 428 - 4692 428 - 4202 411 1965 407 6604 405 3577 395 2439 391 713 375 713 376 7491 368	rotein kinase kinase	7069	558	571
4576 536 2 4093 516 6489 503 7903 500 6872 428 4202 411 1965 407 6604 405 52720 375 713 368 7491 368	(rypsin 1)	6323	558	929
4093 516 6489 503 7903 500 6872 428 4692 428 4202 411 1965 407 6604 405 3577 395 2439 391 713 375 2720 370 7491 368	ed protein	4576	536	2177
6489 503 7903 500 6872 428 4692 428 4202 411 1965 407 6604 405 3577 395 2439 381 713 375 7491 368 4465 366	.18	4093	516	491
7903 500 6872 432 4692 428 4202 411 1965 407 6604 405 3577 395 2439 391 4492 381 713 375 2720 370 7491 368		6489	503	581
6872 432 4692 428 4202 411 1965 407 6604 405 3577 395 2439 391 4492 381 713 375 2720 370 7491 368 4465 366		7903	500	364
4692 428 - 4202 411 1965 407 6604 405 3577 395 2439 391 4492 381 713 375 2720 370 7491 368 4465 366	01.3	6872	432	433
4202 411 1965 407 6604 405 3577 395 2439 391 4492 381 713 375 2720 370 7491 368 4465 366	in H	4692	428 -	96
1965 407 6604 405 3577 395 2439 391 4492 381 713 375 2720 370 7491 368 4465 366		4202	411	381
6604 405 3577 395 2439 391 4492 381 713 375 2720 370 7491 368 4465 366	ating protein 1	1965	407	263
2439 391 4492 381 713 375 2720 370 7491 368 4465 366		6604	405	830
2439 391 4492 381 713 375 2720 370 7491 368 4465 366	smembrane domains	3577	395	431
4492 381 713 375 2720 370 7491 368 4465 366	ed kinase	2439	391	341
713 375 2720 370 7491 368 4465 366	protein 6	4492	381	308
2720 370 7491 368 4465 366	e/glutamine rich -binding protein-	713	375	662
7491 368 4465 366		2720	370	378
4465 366	matrix metalloproteinase 23A	7491	368	138
	n kinase C substrate,	4465	366	335

CO moming contact DO	1649	362	234
	5283	359	507
CCAAT/enhancer binding protein (C/EBP),	7237	359	645
alpha	000	L C	200
paired box gene 9	5206	335	107
protein tyrosine phosphatase, receptor	8235	331	250
type, N		1	
keratin 8	7215	327	449
claudin 7	280	325	173
tronhinin associated protein (tastin)	462	323	360
	8356	322	366
basic helix-loop-helix domain containing,	6734	318	215
class B, 2			C
annexin A2	4467	. 082	977
cathepsin D (lysosomal aspartyl protease)	7370	289	009
Ricandal D (Orosonhila) homolog 1	6822	284	217
Cicatuda D (Diosophina) inclining in	7334	280	310
lectin, galactoside-billollig, soldole, 1 (zalactin 1)	3		
(garconn 1)	7171	278	481
olyceraldehyde-3-phosphate	3953	278	307
giyod aradiyad o priocepriodo			
den you ogon raso an isoporin 6 kidney snecific	7591	277	258
TNE recentor-associated factor 1	6037	269	270
Rho GDP dissociation inhibitor (GDI) alpha	5565	267	321
calcium channel, voltage-dependent,	3142	260	207
gamma subunit 4	2777	VVC	.596
grutamate receptor, fortou opto, kairiate i CCR4-NOT transcription complex, subunit	586 883	244	428
4			
filamin A, alpha (actin-binding protein-280)	3457	241	233
peanut (Drosophila)-like 1	711 .	233	310
enhancer of rudimentary (Drosophila)	5921	231	405
normong endothelin converting enzyme 1	7835	229	197

208 145	164	345	141	116	365	74	98	7	44	201	201	136	108		761	7	5	8/	220	35		141	159	20/	101		Ş	79	700	707		197	
223 217	216	212	207	205	202	200	9	081	190	184	184	178	175	į	174	6 1	172	161	160	158	!	157	155	150	150		1	147	1	14/		144	
2483 7679	4871	2827	7848	2731	999	4489	i	5185	4427	928	928	2111	7852	·	1606		5608	6334	1035	6318		3154	8050	2084	6674			1335	1	2882		1433	1
protocadherin 17	aquapoini o evizancio I	Synapsini Tuhulin alaha brain-specific	CD44 antigen (homing function and Indian	blood group system)	cereptal cavernous manorinamons i	adenylate cyclase activating polypeptide 1	(pituitary)	hairless (mouse) homolog	ujirsuj	oran din	מוביים	graniani postky orowath response 1	cyclin-dependent kinase inhibitor 1A (p21,	Cip1)	stem cell growth factor; lymphocyte	secreted C-type lectin	CD4 antigen (p55)	PCTAIRE protein kinase 1	Rho GTPase activating protein 6	G protein-coupled receptor 37 (endothelin	receptor type B-like)	syndecan 4 (amphiglycan, ryudocan)	PCTAIRE protein kinase 3	empty spiracles (Drosophila) homolog 2	transglutaminase 1 (K polypeptide	epidermal type I, protein-glutamine-gamma-	glutamyltransferase)	potassium voltage-gated channel,	subfamily G, member 2	aldehyde dehydrogenase 4 (glutamate	gamma-semialdehyde dehydrogenase;	pyrroline-5-carboxylare deriyarogeriase/ E14 hinding profein n300	

88	150 92	56	87	91	84	256	29	136	74	91	130	58		C	n t	99	0	8/	139	000	87	. 74	205	09	223	207	1 01	7.5	103
143	134 129	129	128	124	119	118	117	116	114	113	110	109	•		108	105	•	103	101	;	35	95	95	1 00) œ	0 00	/8	84	81
8211	4536 5462	3074	7037	7769	4181	443	1571	2735	9656	5772	6016	4702			8259	3857		2040	2832		2146	7731	1389	 07.7.L	45.59	0280	3641	1874	1783
LIM homeobox transcription factor 1, beta	eukaryotic translation elongation factor 2 mitogen-activated protein kinase kinase	kinase 10 PPAR(gamma) angiopoietin related protein	homen hox A5	CD63 antigen (melanoma 1 antigen)	niclear recentor coactivator 3	CD68 antigen	transmembrane 4 superfamily member 7	nancreatic polyneptide	endothelin type b receptor-like protein 2	neurogenin 1	insulin receptor substrate 2	glutamate receptor, ionotropic, N-methyl D-		binding)	neurotrophin 5 (neurotrophin 4/5)	pyrimidinergic receptor P2Y, G-protein	coupled, 6	empty spiracles (Drosophila) homolog 1	chromobox homolog 2 (Drosophila Pc	class)	heart and neural crest derivatives	expressed 1 transglutaminase 2 (C polypeptide, protein- glutamine-gamma-glutamyltransferase)	(adala) bassis	enolase I, (alpila)	thyroid transcription factor 1	keratin 19	Deleted in split-hand/split-foot 1 region	glutathione peroxidase 4 (phospholipid	hydroperoxidase) aristaless (Drosophila) homeobox

GATA-binding protein 4	629	80	011
transcription factor 1, hepatic; LF-B1,	712	80	136
hepatic nuclear factor (HNF1), albumin			
proximal factor			
heat shock transcription factor 1	6708	79	22
iver-specific bHLH-Zip transcription factor	765	2.2	188
iver-specific bHLH-Zip transcription factor	765	77	188
eukaryotic translation initiation factor 3,	6280	92	94
subunit 4 (delta, 44kD)			1
eukaryotic translation initiation factor 3,	6280	. 92	94
subunit 4 (delta, 44kD)			
gamma-aminobutyric acid (GABA) A	3720	70	178
receptor, alpha 6			
etinoic acid receptor, gamma	5191	70	109
homeo box D9	1776	69	34
MAD1 (mitotic arrest deficient, yeast,	1946	63	88
homolog)-like 1			
homeo box A4	. 7117	22	103
Thy-1 cell surface antigen	559	22	87
	603	53	84
bone morphogenetic protein receptor, type	332	53	45
I (serine/threonine kinase)			;
nepatocyte nuclear factor 3, alpha	72	52	09
vanodine receptor 2 (cardiac)	5244	47	40
signal transducer and activator of	3661	40	32
ranscription 6, interleukin-4 induced			•

APPENDIX 2: Table 6

SwissPrat	Spot	geneName	day3	day8	day	16	aliMean	ratioitoA	rationintto/ classID		className
P04270	E19ab2	actin; alpha; cardiac muscle	1198.174	76	50 59	55,077	2637.676	2.31E+00	-6.57E-01	7	Equal A/Int; Up Islets
O95996	N17ab2	adenomatous polyposis coll like	2708.542	516		94.139		1.49E+00	9.36E-01		Equal Wint: Up Islets
Q14697	G02efB	alpha glucosidase II alpha subun	306.0691 389.822	35 50		5,3358 238.43	.,	1.34E+00 1.67E+00	1.92E-01 3.80E-01		Equal Afint; Up Islets Equal Afint; Up Islets
P14209 P30530	G23ab7 G14ef5	antigen identified by monoclonal AXL receptor tyrosine kinase	314,4639			4.5658		1.01E+00			Equal A/int; Up Islets
043770	123ab3	B-cell CLL/lymphoma 7C	700.573	105	50 1	504.52	1085,931	1.10E+00	5.87E-01		Equal Aint; Up Islets
043852	P08ab3	calumenin	801.582			20.807	2269.921		6.84E-01 -7.52E-01		Equal A/Int; Up Islets
O75718 Q9Y3C0	024cd6 G14el2	cartilage associated protein CGI-116 protein	430.4432 181.0293			090.59 6.7246	592.2327 299.7237	1.34E+00 1.20E+00	7.36E-01		Equal A/Int; Up Islets Equal A/Int; Up Islets
P78369	G09cd6	daudin 10	1176.087			98.595					Equal Alint; Up Islets
P08572	P08ef6	collagen; type IV; alpha 2	1130.23			84.425	2389.488		2.60E-01		Equal A/Int; Up Islets
P11802	B05ef5	cyclin-dependent kinase 4	914.1579			1.1473		1.36E+00 3.01E+00	9.79E-02 5,14E-01		Equal A/Int; Up Islets Equal A/Int; Up Islets
Q18555 P33316	O07ab4 H23ab5	dihydropyrimidinase-like 2 dUTP pyrophosphatase	114,6165 177,2032			1.5382		1.61E+00	4.23E-01		Equal A/Int; Up Islets
P07992	K20ef6	excision repair cross-complemen	362.031		15 90	7.7933	561.6388	1.33E+00	1.97E-01	7	Equal A/Int; Up Islets
075636	B14cd3	ficolin (collagen/fibrinogen domai				27.606		2.00E+00 2.01E+00			Equal A/Int; Up Islets Equal A/Int; Up Islets
Q9UBA6 Q9Y5P5	B19ef4 P08ef7	G8 protein GDP-mannose pyrophosphorytas	370.6842			89,206 4,0862	261.0679		6.41E-01 2.10E-01		Equal Alint, Up Islets
075293	E02ef7	growth arrest and DNA-damage-I				6.6833	261,3841	9.92E-01	6.35E-02	7	Equal A/Int; Up Islets
P50152	C05ab6	guanine nucleotide binding prote	204.743			7.3097		1.28E+00			Equal Afint, Up Islets
Q9NX09	C05gh3 J22ab7	HIF-1 responsive RTP801 high-mobility group (nonhistone (348.8574			32,929 59.895	545,0622 3082.818		-4.61E-01 -8.84E-01		Equal Wint; Up Islets Equal Wint; Up Islets
P05204 Q9P0P2	C17ef3	homolog of yeast MOG1	325.8407			2,7568		1.07E+00			Equal A/Int; Up Islets
	L15gh2	hypothetical protein FLJ10055	172.187	2	41	374.3		1.12E+00			Equal Alint; Up Islets
Q9NWV4	B17gh2	hypothelical protein FLJ20580	221.4432			33.5381 30.9111		1.13E+00 1.11E+00			Equal A/Int; Up Islets Equal A/Int; Up Islets
Q9P0S8 Q9BWS9	A21e/3 P04gh5	hypothetical protein HSPC195 hypothetical protein MGC3234	153.824 196.9154			27.2066		1.12E+00			Equal A/Int; Up Islets
Q9BSK0	K21gh8	hypothetical protein MGC4415	158,4692			34,3425	319.1224		-4.25E-01	7	Equal A/Int; Up Islets
Q9Y6M1	P09cd6	IGF-II mRNA-binding protein 2	135.2566			8.5101		1.59E+00			Equal A/Int; Up Islets
Q9NQX7 P08648	H02gh7	integral membrane protein 3	478.2728			102.244 78.7266	687.2575	1.20E+00 1.3BE+00			Equal A/Int; Up Islets Equal A/Int; Up Islets
O00410	E23ef7 J24ab6	Integrin; alpha 5 (fibronectin rece karyopherin (importin) beta 3	326.6874			29.7656		9.47E-01			
P07195	N16ab6	lactate dehydrogenase B	1003.971	12		593.079	2599.05				Equal Afint; Up Islets
P09382	N22ab6	lectin; galactoside-binding; solub				3086.75 \$3.3883	44205.77	2.03E+00 1.63E+00		7	Equal A/Int; Up Islets Equal A/Int; Up Islets
O75427 Q16553	P20ab6 A15ab7	leucine-rich repeat protein; neuro lymphocyte antigen 6 complex; to				38.4504		1.14E+00			Equal A/Int; Up Islets
075900	005cd4	matrix metalloproteinase 23A	381,2184	2	97 85	51.7221		1.16E+00			
Q9UNF1	H16ef7	melanoma antigen; family D; 2	193.8125	_		59.9701		1.77E+00		7	
P13995 O00265	J16cd6 G19cd8	methylene tetrahydrofolate dehyd microtubule-associated protein; f				59.6021 77.5576	267.156	1.24E+00 9.93E-01			
P27361	B13ef5	mitogen-activated protein kinase				20.3119	486.849				Equal Wint; Up Islets
P24844	E18cd6	myosin; light polypeptide 9; regul				2518.95	11793.09			_	
Q9NVD4	J10gh2	N-acetylneuraminic acid phospha phosphoserine aminotransferase				90,6642 238.851	237.1908 553.0005	1.35E+00 2.87E+00		7	Equal A/Int; Up Islets Equal A/Int; Up Islets
Q9Y617 Q95356	N22ef8 A11gh7	pituitary lumor-transforming 3	237.2126			276.181				-	
Q9Y5X6	G04cd6	plasma glutamate carboxypeptide				66.6202			-2.58E-01	7	
Q15113	G20ab8	procollagen C-endopeptidase en procollagen-proline; 2-oxoglutara				628.947 56,4003		2.40E+00 1.53E+00		7	Equal A/Int; Up Islets Equal A/Int; Up Islets
O15460 O43556	D20cd4 19gh1	sarcoglycan; epsilon	172.719			349.304		1.02E+00			
P50454	M17ab6	serine (or cysteine) proteinase in	339.1762	2 6		091.161		3.59E+00		7	
Q92853	A23cd8	similar to vaccinia virus Hindlil K				97.4551 472.155		1.17E+00 2.49E+00		7	Equal A/Int; Up Islets Equal A/Int; Up Islets
Q16658 Q9Y4Y8	P20cd1 E18cd7	singed-like (fascin homolog; sea Sm protein F	257,2026			48.9705				7	
Q9BVH9	D18gh7	thioredoxin related protein	522.0262		81 19	903.964	968,9954	1.87E+00	-1.18E-01	7	
Q9HBB0	L08gh8	Thy-1 co-transcribed	327.3502			110.527	603.8101	1.76E+00 3.40E+00		7	Equal A/Int; Up Islets Equal A/Int; Up Islets
P01033 Q15582	P09ef7 L20ef6	tissue inhibitor of metalloproteina transforming growth factor; beta-				3029.41 8540.86		2.81E+00		7	
Q01995	M07cd2	Iransgelin	3307.82			8534.27		2.49E+00	2.29E-01	7	Equal A/Int; Up Islets
Q13541	B05ab2	trophoblast glycoprotein	479.648			399.786			-1.70E-01	7	
P06468 P51784	G08cd2 M21cd4	tropomyosin 2 (beta) ublquitin specific protease 11	339,229 158,047			3203.05 91.1246		5.28E+00 1.64E+00		7	
P03996	E17ab2	aclin; alpha 2; smooth muscle; a			40 4	1801.34	14717.63	6.66E+00	2.23E+00	9	
P08123	P02ef6	collagen; type I; alpha 2	186.566	5 5					1.63E+00		Increasing A/Int/I
P24821	A05ef7	hexabrachion (tenascin C; cytota	355.499 374.1490			371.983 474 789			1,37E+00 1,20E+00		Increasing A/Int/I Increasing A/Int/I
Q9NVA2 P40261	F13gh3 M09cd1	hypothetical protein FLJ10849 nicotinamide N-methyltransferas							1.12E+00		Increasing A/Int/I
Q15063	C01ef7	osteoblast specific factor 2 (fasci	276.63	4 7					1.52E+00		Increasing A/InVI
P09486	A09ef7	secreted protein; acidic; cysteine				63837.2 0078.17) 2.77E+00) 1.52E+00		Increasing A/InVI Increasing A/InVI
P35625 Q9H2L5	K20ef5 G06gh8	tissue inhibitor of metalloproteina AD037 protein	216.25 361.229			58.8121		2 -1.19E+00			Int> (A;I)
P15121	C1Bab2	aldo-kelo reductase family 1; me	2342.74	4 53	80 2	337.052	3354.494	-3.51E-03	3 1.20E+00	19	Int> (A;I)
P08582	B14ef5	antigen p97 (melanoma associal							9.92E-01		Int> (A:I)
P20292 O15342	E06ab2 A13ab3	arachidonate 5-lipoxygenase-aci ATPase; H+ Iransporting; lysoso				03.2911 160.831			2.78E+00 2 1.44E+00) int> (A;t)) int> (A;t)
Q9BXJ0	H18gh6	C1q and tumor necrosis factor re			50 1	54.1441	228.0109	-2.26E-01	9.55E-01	19	Int> (A;t)
Q9NRJ3	P09gh4	CC chemokine CCL28	250,002	1 18	30 7	63.2866	948.0274	1.61E+00	2.87E+00	19) Int> (A;I)
P05305	F17ab5	endothelin 1	866.456			403.431 006.711			1.06E+00 1 2.35E+00		9 int> (A;I) 9 int> (A;I)
P09341 Q92730	A04ab6 O14ef4	GRO1 oncogene (melanoma gro GTP-binding prolein	653,857						1 2.08E+00) Int> (A;1)
Q9H5K0	L10gh5	hypothetical protein FLJ23360	151.202		296 1	30.3196	192.6442	2 -2.14E-0	9.71E-01	19) Int> (A;I)
P18564	E21ef7	integrin; beta 6	540.036			33.0055			1.13E+00		3 Int> (A;I)
P14316 P40305	B16ef6 O20ab7	Interferon regulatory factor 2 Interferon; alpha-inducible protein	252.360 t 117.315			55.7047 18.2471			2 9.04E-01 1 2.71E+00		9 Int> (A;I) 9 Int> (A;I)
P05161	B21cd5	interferon-stimulated protein; 15				26.6346			1 1.70E+00) Inl> (A;I)

D40704	100.5								
P42701	A06ef7	Interleukin 12 receptor; beta 1	1227.347	22100	1333,521		1.20E-01		19 Int> (A;I)
P09237	J15ef1	matrix metalloproteinase 7 (matri		87300	19849.69		1.05E+00		19 lni> (A;i)
P16860	O13ab8	natriuretic peptide precursor B	4340.214	12900	1260,906		-1.78E+00		19 Int> (A;I)
P21359	L16ab7	neurofibromin 1 (neurofibromatos	132.483	292	155.0822	193.3208	2.27E-01		19 Int> (A;I)
P02775	B07ef7	pro-platelet basic protein (include		1400	265.2119		2.96E+00		19 Int> (A;I)
Q9UDQ9		SBBI26 protein	121.5886	1080		479.5908			19 Ini> (A;I)
075635	P14cd3	serine (or cystelne) proteinase in		3890	326.2581		4.88E-01		19 Int> (A:1)
P02735	A11cd3	serum amyloid A1	496.9578	17400	957.6141	6295.724	9.46E-01		19 Int> (A;I)
060635	F10cd5	tetraspan 1	321.8888	843	177.6479		-8.58E-01		19 thi> (A;i)
P50591	C09cd4	tumor necrosis factor (ligand) sur		1550	476.2305		-1.73E-01		19 Int> (A;I)
Q03169	110ef7	lumor necrosis factor; alpha-indu		1260	326,6536		-1.67E-01		19 int> (A;i)
Q9C075	N11ef8	type I intermediate filament cytok		4200	401.4854		-1.08E+00		19 int> (A;i)
P02248	A01cd3	ubiquitin C	155.6735		62.04595		-1.33E+00		19 int> (A;I)
O95497	A15cd5	vanin 1	309.1556	1840		887.0948			19 Int> (A;I)
Q14202	B15cd5	zinc finger prolein 261	199.6246	1040	247.9161	496,3668	3.13E-01		19 Int> (A;I)
P15514	M02ef7	amphiregulin (schwannoma-deriv	103.1419	405	65,66067		-6.52E-01		18 Off A/I; On Int
Q9BZL9	121 gh8	B aggressive lymphoma gene	126.3951	211	84.56927		-5.80E-01		18 Off A/I; On Int
P30991	D14ef5	chemakine (C-X-C motif); recepts		260	120,2328	148.3131	9.01E-01	2.02E+00	18 Off A/I; On Int
P29400	H13ab3	collagen; type IV; alpha 5 (Alport		374	141.1609	189.6904	1.40E+00		18 Off A/I; On Int
Q9UK22	H22cd7	F-box only protein 2	87.59822	217	120.3781	141.6466	4.59E-01		18 Off A/I; On Ini
Q16769	O15cd8	glutaminyl-peptide cyclotransfera		195	113,0683	132.7502	3.31E-01		18 Off A/I; On Int
P02261	M06gh6	H2A histone family; member I	65.46028	263	126.664	151.868	9.52E-01	2.01E+00	18 Off A/I; On Int
P20769	D03ef1	Immunoglobulin heavy constant r	101.245	176	102.5687	126,7136	1.87E-02	8.00E-01	18 Off A/I; On Int
P52945	112ab6	insulin promoter factor 1; homeox		351	86.42558	170.7555	2.12E-01		18 Off A/1; On Int
Q14496	F21cd6	Interferon-induced protein 44	99.27101	286	98.28057	161.3017	-1.45E-02	1.53E+00	18 Off A/I; On Int
P01583	F15ef7	Interleukin 1; alpha	103,386	375	66.6314	181.5566	-6,34E-01	1.86E+00	18 Off A/I; On Int
P01584	F17ef7	Interleukin 1; beta	116.7675	415	119.0156	216.8707	2.75E-02	1.83E+00	18 Off A/I; On Int
Q9GZM1	120gh7	NDRG family member 4	102.1328	348	101.569		-7.99E-03		18 Off A/I; On Int
P25105	G15ef5	platelel-activating factor receptor		356		177,9689			18 Off A/I; On Int
P58294	O14gh8	prokinelicin 1 precursor	79.98286	241	137.5167	152,9437	7.82E-01	1.59E+00	18 Off A/I; On Int
P23471	A06ef6	protein tyrosine phosphatase; rei	85,55935	346	93.65845	175.0253	1.30E-01	2.02E+00	18 Off A/I; On Int
O88386	D10ef3	RAB10; member RAS oncogene	123,2003	399	68.13944	196.7342	-8.54E-01	1.69E+00	18 Off A/I; On Int
P10826	108gh6	retinoic acid receptor; beta	67.98871	241	120.9469	143.3844	8.31E-01	1.83E+00	18 Off A/I; On Int
O95786	124ef3	RNA helicase	112,9356	241	89.32396	147.8829	-3.38E-01	1.10E+00	18 Off A/I; On Int
P02778	O10ef7	small inducible cytokine subfamil	65.03395	418	86.09019	189.6022	4.05E-01	2.68E+00	18 Off A/I; On Int
P80162	F14cd1	small inducible cytokine subfamil	54,00821	311	129.2639	164.6594	1.26E+00	2.52E+00	18 Off A/I; On Int
P43005	N23ef5	solute carrier family 1 (neuronal/	99.30514	189	110.8119	133,1898	1.58E-01	9.32E-01	18 Off A/I; On Int
P01375	O09ef6	tumor necrosis factor (TNF super	138,6153	386	75.13436	200.0804	-8.84E-01	1.48E+00	18 Off A/I; On Int
Q16890	G04cd2	tumor protein D52-like 1	103,5477	191	98.01862		-7.92E-02	8.84E-01	18 Off A/I; On Int
Q9H949	K11gh5	WW45 protein	111,8812	218	79.54574	136,5769	-4.92E-01	9.64E-01	18 Off A/I; On Int
Q9H4G4	O06gh5	17kD fetal brain protein	88.61746	223	532.2979	281.2148	2.59E+00	1.33E+00	8 Off Adinar; Increasing Int/I
Q14040	B15ef1	collagen; type VI; alpha 1	38.27203	209	4724.413	1657.194	6.95E+00	2.45E+00	8 Off Adnar, Increasing Int/I
P12110	B17ef1	collagen; type VI; alpha 2	88.00282	164	3835,637	1362.487	5,45E+00	8,97E-01	8 Off Acinar; Increasing Int/I
P02751	K15ef1	fibronectin 1	4.441362	152	4171.603	1442.812	9.88E+00	5.10E+00	8 Off Acinar; Increasing Int/I
P14652	N23ab6	homeo box B2	62.71629	291	3571,974	1308.575	5.83E+00	2.21E+00	8 Off Acinar, Increasing Int/I
P08476	D15ef7	inhibin; beta A (activin A; activin	64.38161	318	854.6532		3.73E+00		8 Off Acinar, Increasing Int/1
P17936	D07ab6	insulin-like growth factor blnding	38.22268	349	1302.669	563.1937	5.09E+00	3.19E+00	8 Off Acinar, Increasing Int/I
Q9Y4K0	P18ab6	lysyl oxidase-like 2	75,91549	260	2525,644	953.7036	5.06E+00	1.77E+00	8 Off Acinar, Increasing Int/I
P03956	N05ef7	matrix metalloproteinase 1 (inter:	106,2193	203	590.2622	299.8442	2.47E+00	9.35E-01	8 Off Acinar, Increasing Int/I
P39900	N15ef7	matrix metalloproteinase 12 (mac	18.2357	381	1219.953		6.06E+00		8 Off Adinar, Increasing Int/I
P55001	G09ab7	microfibrillar-associated protein 2	99.33548	526	3302,631	1309,293	5.06E+00	2.40E+00	8 Off Acinar, Increasing Int/I
Q9NYR0	007gh7	SH3-domain kinase binding prote		197	656.4675	296,0285	4.24E+00	2.50E+00	8 Off Acinar, Increasing Int/I
043623	D06ef4	snail hemolog 2 (Drosophila)	96.88079	292	3028,508	1139.155	4.97E+00	1.59E+00	8 Off Acinar; Increasing Int/I
P09936	113cd4	ubiquilin carboxyl-terminal estera	48.30052	219	1523.166		4.98E+00		8 Off Acinar; Increasing Int/I
Q02952	J11ab2	A kinase (PRKA) anchor protein	103.9756	364	229.1446	232.3779	1.14E+00	1.B1E+00	4 Off Adnar; On Int=I
Q99541	119ab2	adipose differentiation-related pr		182	146.1347		1.12E+00		4 Off Acinar; On Int=I
P54284	N17ab3	calcium channel; voltage-depend		227	254.9633		1.26E+00		4 Off Acinar; On Int=1
Q13269	G14ab5	cAMP responsive element bindin		193	217.7521		1.11E+00	9.40E-01	4 Off Acinar; On Int≈I
Q9NPF2	M02ef4	chondroitin 4-sulfotransferase	133.8375	192	303.01		1.18E+00		4 Off Acinar, On Int=1
Q9H7A5	F12gh7	chromosome 20 open reading fra		179	208.0134		1.08E+00	8,58E-01	4 Off Adnar, On Int=I
Q9H9Q2	L23gh5	COP9 constitutive photomorphos		115	184.3088		1.09E+00	4.14E-01	4 Off Acinar; On Int=1
P42771	D17ef5	cyclin-dependent kinase inhibitor		471		290.1928			4 Off Acinar, On Int≃l
Q14650	J20ef4	cytoplasmic FMR1 interacting pro		272		242.2271			4 Off Acinar; On Int=1
Q961K6	F14gh6	DKFZP434C245 protein	145.227	104	91.06477		-6.73E-01		4 Off Acinar; On Int≈I
P12034	D05ef7	fibroblast growth factor 5	60.2377	259		212,2747			4 Off Adnar; On Int≔I
Q93079	M20gh6	H2B histone family; member J	144.6906	428		281.3281			4 Off Adnar; On Int=1
Q9P016	A10ef8	HSPC144 protein	123.4601	348	399.4935		1.69E+00		4 Off Acinar; On Int≃I
Q9HBI5	M09gh4	HT021	116.3605	276	309.5798		1.41E+00		4 Off Acinar, On Int=1
Q9BY45	O24gh8	HTPAP protein	78.49388	109	177.2043		1.17E+00		4 Off Adnar; On Int=1
P08397	G18ab6	hydroxymethylbilane synthase	139.0857	168		196.1361			4 Off Adnar, On Int=1
Q9BUV0	M06gh4 N22gh6	hypothetical protein dJ465N24.2.		152		129.3473			
Q9H9A2		hypothetical protein DKFZp762U		110	169.5379		1.04E+00		4 Off Acinar; On Int=I
Q9NUV6	P09gh3	hypothetical protein FLJ11113	87.43713	178	136.6771		6.44E-01		4 Off Acinar; On Int=I
Q9NXF7	C20gh2	hypothetical protein FLJ20280	66.49705	143		126.7851			4 Off Acinar; On Int=1
Q9H6V0	M05gh5	hypothetical protein FLJ21839	116.3704	258		232.5844			4 Off Actnar; On Int=1
		hypothetical protein MGC4707	80.31127	170		151.3887			4 Off Acinar; On Int=1
Q9H765	115gh6	hypothetical protein MGC5540	106.3949	220		177.6046			4 Off Adnar; On Int=i
Q14157	N23gh1	KIAA0144 gene product	95.13957	147		129.9305			4 Off Acinar, On Int=1
Q9Y2D8	A07ef8	KIAA0923 protein	84,1225	237	206.8269		1.30E+00		4 Off Acinar; On Int=1
Q9H6Z3 Q99748		kinesin family member 13A	111,6382	206	238.8822		1.10E+00		4 Off Adnar; On Int=1
	E24gh5	pourturdo					4 44F-01	2.74E-01	4 Off Acinar; On Int=1
	J24ab8	neurlurin	123.7486	150					
Q9NVD7	J24ab8 D11gh3	parvin; alpha	82.67948	140	230.66	151,1753	1.48E+00	7.62E-01	4 Off Adnar; On Int=I
Q9NVD7 Q43175	J24ab8 D11gh3 H24cd6	parvin; alpha phosphoglycerate dehydrogenas	82.67948 98.81456	140 119	230.66 192.3326	151,1753 136.6151	1.48E+00 9.61E-01	7.62E-01 2.64E-01	4 Off Adnar; On Int=I 4 Off Adnar; On Int=I
Q9NVD7 Q43175 Q9Y253	J24ab8 D11gh3 H24cd6 E24cd1	parvin; alpha phosphoglycerate dehydrogenas polymerase (DNA directed); eta	82.67948 98.81456 108.2715	140 119 101	230.66 192.3326 157,8762	151,1753 136.6151 122.2465	1.48E+00 9.61E-01 5.44E-01	7.62E-01 2.64E-01 -1.06E-01	4 Off Adnar; On Int=1 4 Off Adnar; On Int=1 4 Off Adnar; On Int=1
Q9NVD7 Q43175	J24ab8 D11gh3 H24cd6	parvin; alpha phosphoglycerate dehydrogenas	82.67948 98.81456 108.2715 87.66238	140 119 101 280	230.66 192.3326 157.8762 172.8525	151,1753 136.6151 122.2465	1.48E+00 9.61E-01 5.44E-01 9.80E-01	7.62E-01 2.64E-01 -1.06E-01 1.67E+00	4 Off Adnar; On Int=I 4 Off Adnar; On Int=I

	13636	K19cd7	RAB31; member RAS oncogene	76,36263	118	170.0352	121.5169 1.15E+00	6.30E-01	4 Off Adinar, On Int=1
		L10ef2		80.78879	171	194,395	148.7422 1.27E+00	1.08E+00	4 Off Adnar; On int=1
	9NVQ7	K12gh3	Sec61 alpha form 2	82.6036	143	162.0674	129.1434 9.72E-01		4 Off Adnar, On Int=1
	75368	J08cd1	SH3 domain blinding glutamic act		212	196.8753	167.1823 1.08E+00		4 Off Adnar, On Int=1
C	95863	N01cd2		76.58448	134	157.9807	122.817 1.04E+00		4 Off Adnar; On Int=1
	29UL01	H01cd8	squamous cell carcinoma antiger		111	176.4557	124.5829 1.03E+00 364.6675 2.25E+00		4 Off Adnar; On Int=I 4 Off Adnar; On Int=I
	16226	N05cd6		111.1384 79,00174	454 139	529.2322 163.2593	127.0819 1.05E+00		4 Off Acinar; On Int=1
	07996	A11ef7	thrombospondin 1 transcription factor 12 (HTF4; he		181	256.0348	182.9347 1.19E+00		4 Off Acinar; On Int=1
	199081 195922	F10ef6 L06ef7		73,12945	245	253.9213	190.8311 1.80E+00		4 Off Acinar, On Int=I
	15036	H18ef6	v-ets erythrobiastosis virus E26 (114	83.55955	115.8531 -8.43E-01	-3.94E-01	4 Off Adnar; On Int=1
	95337	D04ef8	weakly similar to glutathione perc	69,55111	164	284.3593	172.7489 2.03E+00	1.24E+00	4 Off Acinar; On Int=1
	29H1B5	M18gh5	xylosyltransferase II	81,87715	106	174.2539	120.5878 1.09E+00		4 Off Acinar, On Int=I
F	55263	K07ab2	adenosine kinase	891,6698	720	299.6337	637.198 -1.57E+00		15 On A/Int; Down I
	51648	A22ab2	aldehyde dehydrogenase 3 famil		1390	507.3786	1284,046 -1.95E+00		15 On Affait Down i
	50995	108ab2	annexin A11	4888.71	3870	1773.94	3511.843 -1.46E+00 7424.351 -2.50E+00		15 On A/Int; Down I 15 On A/Int; Down I
	212429	114ab2	annexin A3	9711.235	10800 5290	1720.452 1976.367	3744.962 -1.00E+00		15 On A/Int; Down I
	205026	L22ab2 M07ab3	ATPase; Na+/K+ transporting; b∈ B-factor; properdin	2984.309	2460	1178.473	2206.568 -1.34E+00		15 On A/Int; Down I
	P00751 Q9HA23	E10gh6	calponin like transmembrane dor		528	193.0456	400.5912 -1.32E+00		15 On A/Int; Down I
	216170	M17ab3	carcinoembryonic antigen-relater		1230	212.7712	807.5766 -2.21E+00		15 On A/Int; Down I
	P25774	P07ab5	cathepsin S	857,9751	776	228.9093	620.9386 -1.91E+00	-1,45E-01	15 On A/Int; Down 1
	P21926	J13ef7	CD9 antigen (p24)	2011.966	3500	833.332	2115.74 -1.27E+00		15 On A/Int; Down I
(Q9H2A7	M19gh5	chemokine (C-X-C motif) ligand 1		842		576.1228 -1.46E+00		15 On A/Int; Down !
	014493	G21ab4	claudin 4	1919.128	2480	799.5104	1732.749 -1.26E+00		15 On A/Int; Down I 15 On A/Int; Down I
	P12277	M06ab4	creatine kinase; brain	373,8119	617	193.0681	394,4895 -9.53E-0		15 On A/Int; Down I
	Q9NYS7	E20ef4	CS box-containing WD protein	596,6891	384 1260	211,1821 558,9758	397.3278 -1.50E+00		15 On A/Int; Down I
	P53355	D09ab5	death-associated protein kinase	1325.243 7363.301	14700	2196.358			15 On Alint; Down I
	015205	D17cd6	diubiquitin DKFZP564K247 protein	1325.797	1060	313,5237	900,9109 -2.08E+0		15 On A/Int; Down I
	Q9UFZ2 P28562	C23ef8 M22ab5	dual specificity phosphalase 1	2077.188	1680				15 On A/Int; Down I
	Q16690	B21ab4	dual specificity phosphatase 5	485.6475	305	155.0161			15 On A/Int; Down I
	Q9UJW0	H10ef1	dynaclin 4 (p62)	778.5079	662	335,9831	592.1023 -1.21E+0	-2.34E-01	15 On A/Int; Down I
		E13gh5	ectonucleotide pyrophosphatase.		378	183,3797			15 On A/Int; Down I
- 1	P00533	A06ef5	epidermal growth factor receptor	263.2516	334	182,4044			15 On A/Int; Down
- (Q13113	J02cd5	epithelial protein up-regulated in	13324,19	15500	2641,961	10494.6 -2,33E+0		15 On A/Int; Down I
	P28161	M09ab6	glutathione S-transferase M2 (ma		690	217.4433			15 On A/Int; Down I
	Q03013	L06ab3	glutathione S-transferase M4	333.2767	465	210.8696			15 On A/Int; Down I 15 On A/Int; Down I
	P52594	A18ab7	HIV-1 Rev binding protein	439,4544	468 757	252.8058 368.3558			15 On A/Int: Down I
	Q9NWT9		hypothetical protein FLJ20607 hypothetical protein HT036	1062.388 748.0681	844	160,1449			15 On Alint; Down I
	Q9BZR4 Q9BW25	L12gh7 A19gh6	hypothetical protein MGC3101	315.4695	409	150.4768			15 On A/Int; Down I
	Q96W25	I16ab6	integrin; beta 4	490.7243	672	157,31			15 On A/Int; Down I
	P10145	O22ef7	Interleukin 8	6902.736	13300				15 On A/Ini; Down i
	Q05084	G10ab7	islet cell autoantigen 1 (69kD)	600.5671	588	209.1329			15 On A/Int; Down I
	Q9Y484	E14cd7	JM5 protein	1335.218	2010				15 On A/Int; Down I
	P14923	M23ef6	junction plakoglobin	1969,949	2610				15 On A/Int; Down I
	P05783	N06ef6	keratin 18	17317.24	12500				15 On A/Int; Down I 15 On A/Int; Down I
	P08727	L06ab6	keratin 19	4025.258	2800				15 On A/Int; Down I
	P08729	N14ef6	keratin 7	11605.48 11031.01	11100 7880	1888.38 1694.024			15 On A/Int; Down I
	P05787	N16ef6 H22ab6	keralin 8 kinesin family member 3C	366,3528	652				15 On A/Int; Down I
	O14782 O00515	D05ab7	ladinin 1	3562.632	2040				15 On A/Inl; Down I
	P80188	D13ab7	lipocalin 2 (oncogene 24p3)	5234.435	9070				15 On A/Int; Down I
	Q9UQ53	G02ef3	mannosyl (alpha-1;3-)-glycoprote		931				15 On A/Int; Down I
	P80294	D15gh6	metallothionein 1H	327.8489	366			1 1.57E-01	15 On A/Int; Down I
	075394	M10cd5	mitochondrial ribosomal protein l	441.6574	646				15 On A/Int; Down I
	Q9Y376	H18ef2	MO25 protein	860.3225	1410				15 On A/Int; Down I
	P15941	H22ab7	mucin 1; transmembrane	2330.649	1250				15 On A/Int; Down I 15 On A/Int; Down I
	Q16301	H11cd4	myelin transcription factor 2	2433.02	2560 1110				15 On A/Int; Down I
	Q99836	M07ab7 H08cd4	myeloid differentiation primary re natural killer cell transcript 4	49232.84	71300				15 On A/Int; Down I
	P24001 Q9C002	O12gh8	normal mucosa of esophagus sp		17700				15 On A/Int; Down I
	P25963	A16ef1	nuclear factor of kappa light poly		1270				15 On A/Int; Down I
	P00491	L18ab7	nucleoside phosphorylase	820.4015	500				15 On A/Int; Down I
	P49763	Q20ef7	placental growth factor; vascular	5604.053	12600	1723.953	6646.258 -1.70E+0		15 On A/Int; Down I
	Q9Y342	A13ef2	plasmolipin	942.5595	983				15 On A/Int; Down J
	P43490	H04cd5	pre-B-cell colony-enhanding fact		1730				15 On A/Int; Down I
	Q99988	B23ef7	prostate differentiation factor	1470.061	2930				15 On A/Int; Down I 15 On A/Int; Down I
	Q9UJY1	B05ef3	protein kinase H11	1836.927	2080		1 1485.828 -1.75E+(15 On Allnl; Down I
	Q9H1C7	O10gh8	putative nuclear protein ORF1-F		14800 5250		9771.26 -1.90E+0 7 3646.784 -5.94E-0		15 On A/Int; Down I
	P51149	A21ef6 J22cd4	RAB7; member RAS oncogene f RAB9A; member RAS oncogene				1393.904 -1.38E+		15 On A/Int; Down I
	P51151 P15153	C17ef6	ras-related C3 botulinum toxin si	383,0769	418				15 On A/Int; Down I
	P52566	119ef1	Rho GDP dissociation inhibitor ((986.1501	788		708.6827 -1.49E+		15 On A/Int; Down I
	P01011	108ef7	serine (or cysteine) proteinase in		43700				15 On A/Int; Down I
	P05120	, P07ef7	serine (or cysteine) proteinase it		38200		9 21999.69 -3.11E+	00 6.16E-01	15 On A/Int; Down I
	014508	M13ef6	STAT induced STAT inhibitor-2	1511.844	1490				15 On A/Int; Down I
	Q9Y6N5		sulfide dehydrogenase like (yea		871		7 659,5107 -1.54E+		15 On A/Int; Down I
	000161	C23cd4	synaptosomal-associated protein		1340		2 1034.031 -6.50E-		15 On A/Int: Down I
	P31431	F24cd1	syndecan 4 (amphiglycan; ryudo		3080				15 On A/Int; Down I 15 On A/Int; Down I
	015533	B16ef7	TAP binding protein (tapasin)	313.1585	424		1 299.8834 -9.48E-		15 On A/Int; Down I
	P17987	E11ef5	t-complex 1	495.6787	298 373				15 On A/Int; Down I
	014748	O20ef6 C06ef1	telomerase reverse transcriptase transmembrane 4 superfamily m		1740				
	P30408	E07gh5	transmembrane protein 8 (five n				5 806.2983 -1.04E+		
	MALION?		interior process o gave to				The state of the s		

Q16149	B15cd1	transporter 1; ATP-binding casse	862.1561	1100	480.8685	813.3307 -8.42E-01	3.47E-01	15 On A/Int; Down I
Q9Y2A9	C22ef3	UDP-GlcNAc:belaGal bela-1;3-N		436	153,9485	454.8446 -2.33E+00		15 On A/Int; Down I
O60625	N10cd3	veside-associated membrane pn		2680	584.39		1.10E-01	15 On A/Int; Down I
P07948	A21ab7	v-yes-1 Yamaguchi sarcoma vira	512.646	443	170.1742	375,1466 -1,59E+00		15 On A/Int; Down I
Q9P2N4	M10gh4	a disintegrin-like and metalloprot		227	72.96867	153.1495 -1.13E+00		14 On A/Int; Off I
P47895	C04ab2	aldehyde dehydrogenase 1 famili		2370	146,6644	1254.954 -3.09E+00	9.27E-01	14 On A/Int; Off I
P04233	G09ef1	CD74 antigen (invariant polypept		174	86.99566	163,6002 -1,40E+00		14 On A/Int; Off I
Q9NZ31	L12gh4	chromosome 20 open reading fra		375	137,4415		1.18E-01	14 On A/Int; Off I
O96002	G07cd5	chromosome X open reading fran	401.521	375	130.7318	302.4558 -1.62E+00		14 On A/Int; Off I
O94907	J15gh6	dickkopf homolog 1 (Xenopus lae		200	84.53571	145.044 -8.30E-01		14 On A/Int; Off I
Q92796	K06ef5	discs; large (Drosophila) homolo		154	84.90819	133.5184 -9.32E-01		14 On A/Int; Off I
Q13115	017ab4	dual specificity phosphalase 4	533.5415	730	146.312	469.8392 -1.87E+00		14 On A/Int; Off I
P29317	F02ef5	EphA2	267.722	279	82.15254	209.5143 -1.70E+00		14 On Alint; Off I
P29323	D10ef5	EphB2	269,1596	232	74.35016	191.9946 -1.86E+00		14 On A/Int; Off)
Q9UKF9	H10cd7	els hornologous factor	603.0262	587	102.0333	430.6372 -2.56E+00		14 On A/Int: Off I
Q9NPD3	P22gh2	-		300	49.72784	207.9117 -2.46E+00		14 On A/Int; Off I
Q3147 D3	G12ab4	exosome component Rrp41 forkhead box O3A	273.527	169	102.6435	177.9385 -1.35E+00		
P21217	M16ab3	fucosyltransferase 3 (galactoside	262.2734	257	60.24803			14 On A/Int; Off I 14 On A/Int; Off I
075205	D24ef2			382	90.93956	168.8097 -1,65E+00		14 On A/Int: Off I
075712	A03ef1	G protein-coupled receptor; famil				364.8596 -2.77E+00		
		gap junction protein; beta 3; 31kl		402	145.5466	260.8655 -6.93E-01		14 On Afint; Off I
Q92908	D12ab4	GATA binding protein 6	293.117	199	93.59788	195.3976 -1.65E+00		14 On A/Int; Off I
O95210 O95395	P05cd4	genethonin 1	331.9855	241	112.69	228.6851 -1.56E+00		14 On A/Int; Off i
	K21cd5	glucosaminyl (N-acetyl) transfera		274		205.5863 -2.70E+00		14 On A/Int; Off I
P48506	F22ab5	glutamate-cysteine ligase; cataly		243	134.9982	225.879 -1.15E+00		14 On A/Int; Off I
Q9UI98	F08ef8	hqp0256 protein	357.587	336	106.803			14 On A/Int; Off I
Q9H6D8	D18gh5	hypothetical protein FLJ22362	542.6184	595	127.4299	421.5312 -2.09E+00	1.32E-01	14 On A/Int; Off I
P05362	102ab6	intercellular adhesion molecule 1		424	67.80016	301.0074 -2.60E+00	4.56E-02	14 On A/Int; Off I
Q07627	J20gh7	keratin associated protein 1.1	288.3566	303	40.28571	210.3957 -2.84E+00		14 On A/Int; Off I
Q9BYQ7	D10gh8	keratin associated protein 4.10	493.4934	605	96.51137	398.2261 -2.35E+00	2.93E-01	14 On A/Int; Off I
075071	F15gh1	KIAA0494 gene product	240.2512	167	91.77931	166.3595 -1.39E+00		14 On A/Int; Off I
Q9UPQ2	G22ef8	KIAA1100 protein	224.7248	349	135.6811	236.4102 -7.28E-01	6.34E-01	14 On A/Int; Off I
O43896	H10cd6	kinesin family member 1C	159.6982	170	73.44924	134.4385 -1.12E+00		14 On A/Int; Off I
Q13887	L05ab5	Kruppel-like factor 5 (intestinat)	494.1259	376	97.43054	322.6178 -2.34E+00		14 On A/Int; Off I
000312	D22cd3	MAP kinase-interacting serine/thi		150	86.60695	128,2701 -7.73E-01		14 On A/Int; Off I
Q15264	H07ab8	milogen-activated protein kinase		160	76.32069	150.0667 -1.49E+00		14 On A/Int; Off I
Q9UHA4	P10gh1	mitogen-activated protein kinase		244	72.50042	195.0133 -1.89E+00		14 On A/Int; Off I
P21397	F23ef6	monoamine oxidase A	427.3495	480	30.42645	312.5207 -3.81E+00	1.67E-01	14 On A/Int; Off I
P55196	N07ab7	myelold/lymphoid or mixed-linear		168	78.62952		2.56E-02	14 On A/Int; Off I
Q12965	124ab7	myosin IE	212.4935	276	122.0015		3.75E-01	14 On A/Int; Off I
Q9HBW1	O10gh7	NAG14 protein	417.9837	396	86.84398			14 On A/Int; Off I
Q9H2W4	D20gh6	neural precursor cell expressed;		463	75.41197			14 On A/Int; Off I
Q9NRR3	O06gh4	non-kinase Cdc42 effector protei		327	117.0936			14 On A/Int; Off I
P23511	M19ab8	nuclear transcription factor Y; alp		226	132.3351	199,2783 -8.54E-01		14 On A/Int; Off I
Q99650	D14cd4	oncostalin M receptor	337.1614	233	141.0848	237.114 -1,26E+00		14 On Alint; Off I
P04085	016ef7	platelet-derived growth factor alp		463	124.4793	343.2015 -1.83E+00		14 On A/Int; Off I
Q13048 P10586	N06cd2	pregnancy specific beta-1-glycop		203	84.53571	145.8869 -8.25E-01		14 On A/Int; Off I
	E11ef6	protein tyrosine phosphatase; rea		405	136.6718	335.8682 -1.77E+00		14 On Allnt; Off I
O95200	N22ab8	relinoic add receptor responder		315	72.62859			14 On Alint; Off I
Q9NVX8 Q15418	P17ef3 N14ef5	Rho GTPase activating protein 8		479 361	109.2541	337.8544 -1.96E+00		14 On A/Int; Off I
Q15418	F10ab8	ribosomal protein S6 kinase; 90k			78.38666			14 On A/Int; Off I
Q15434	O10cd6	RNA binding motif; single strands		328 235	145.2438			14 On A/Int; Off I
O60679		Sec23 homolog B (S. cerevisiae)			126,9582			14 On Allnt; Off I
Q12B90	D15ef5	serum-inducible kinase	202.0368	194	62.20916			14 On A/Int; Off I
Q12971	N06ef5	SFRS protein kinase 1	350.4057	278	113.0428			14 On A/Int; Off I
P78556	L03cd6 M02cd2	sialyltransferase	247.3141	329 382	96.91002			14 On A/Int; Off I
075751	F14ef4	small inducible cytokine subfamil solute carrier family 22 (extraneu			59.31325			14 On A/Int; Off I
Q9UM01	D08cd4			238	66.83593			14 On A/Int; Off I
		solute carrier family 7 (cationic as		260	55,89528			14 On Affair Off I
Q9Y5X1 P08842	D10ef2 D19ab3	sorting nexin 9 steroid sulfatase (microsomal); a	219.1839	240	108.9101			14 On A/Int; Off I
O43760	G09cd5		391,0545	414 238	136.2997	328,904 -1,68E+00		14 On A/Int; Off I
075674	N15cd5	synaptogyrin 2	276,5234	236 285	118.1167 54.32353			14 On A/Int; Off I
P01135	K04ef7	target of myb1-like 1 (chicken) transforming growth factor; alpha		160				14 On Allnt; Off I
Q9NT70	K13gh3	transmembrane protein vezatin	365.9385	401	59.63082			14 On A/Int; Off I
O60656	G13gh3	UDP glycosyltransferase 1 family		438	129.2607			14 On A/Int; Off I
						337.3961 -4.31E+00		14 On A/Int; Off I
Q9NZ42 O76080	N18gh4 E14cd4	uncharacterized hematopoietic st zinc finger protein 216	314.9706	213	115.2737			14 On A/Int; Off I 14 On A/Int; Off I
Q13015	H11ab2	ALL1-fused gene from chromoso		211 1730	1752.9	200.0768 -2.08E+00		
P04083	106ab2		1114.486			1272.138 2.41E+00 5883.787 2.90E+00		6 On Acinar; Up Int=1
O60592	B14ab2	annexin A1		3320				6 On Adinar, Up Int=I
Q07814	E04ef6	Arg/Abl-interacting protein ArgBF BCL2-associated X protein	430,8903		1101.457	2304.427 3.56E-01 797.9572 1.35E+00		6 On Adinar, Up Int=1
Q14201	E17cd7	BTG family; member 3	197.8608	862 451		355.9496 1.08E+00		6 On Adnar; Up int=1 6 On Adnar; Up int=1
Q9NR00	J21gh4	chromosome 8 open reading fran		7310				6 On Adnar; Up Int=1
Q9BUW7		chromosome 9 open reading fran		650		4453.955 2,47E-01 476.5934 1.27E+00		
095401	N04cd4	cofactor required for Sp1 transcri		395	588.6133			6 On Acinar; Up Int≂l 6 On Acinar; Up Int=l
Q00535	B09ef5	cyclin-dependent kinase 5	269.7495	395 476	432.1176			6 On Acinar; Up Int=1
P21741	E17ab7	midkine (neurite growth-promotin		15400		9388.855 1,71E+00		
Q9Y291	J15ef2	mitochondrial ribosomal protein 5		330	365.4484			6 On Adnar; Up Int=1
075376	E17cd6	nuclear receptor co-repressor 1	375.3679	1270		290,8529 1,04E+00 791,9381 9,54E-01		6 On Acinar; Up Int=1 6 On Acinar; Up Int=1
P03973	N10cd1	secretory leukocyte protease inhi		2070	4153,51			6 On Acinar; Up Int=1
015427	F06cd4	solute carrier family 16 (monocar				589.0175 1.78E+00		6 On Adnar; Up Int=1
P16949	D11ab7	stalturin 1/oncoprotein 18	728.2643	2630		1781.654 1.45E+00		6 On Adnar, Up Int=1
Q9NZ86	J08gh3	uncharacterized bone marrow pri			536,938			6 On Acinar; Up Int=1
000748	H07cd4	carboxylesterase 2 (Intestine; livi		220				20 Other
Q13630	M01cd3	tissue specific transplantation an	622,3905	287	255.576			20 Other
P00505	C14ab6	glutamic-oxaloacetic transaminas			260.2263			20 Other

P50440	A09ab5	glycine amidinotransferase (L-arı	1737.906	39	5	261.2318	798.1958				Other
P29622	A06cd1	serine (or cystelne) proleinase in	461,3518	18	9	264.9574		-8.00E-01			Other
O14646	C17ab4	chromodomain heilcase DNA bin		18		273.1703		1.67E+00			Other
P48307	110cd4	tissue factor pathway Inhibitor 2	920.3007	36		277.6619		-1.73E+00 ·			Other Other
Q15125	D08cd6	emopamil binding protein (sterol	897.4306	23		282.4317	211.5128	-1.67E+00 - 5.26E-01			Other
P19387	B13ab8	polymerase (RNA) II (DNA direct	177.983	15 27		282.8549 283.8007		6.73E-01			Other
. Q9P0K7	H08gh6	retinoic acid induced 14	634.2686	46		300.6211		-1.08E+00			Other
P30086	E04ab8 C06ab5	prostatic binding protein flightless I homolog (Drosophila)		50		306,0777		-1.02E+00			Other
Q13045 O95084	G10cd7	prolease; serine; 23	1096.756	24		314.746		-1.80E+00		20	Other
P49770	L11ef4	eukaryotic translation initiation fa		54	7	329,7361		-1.15E+00			Other
Q13948	J02ef6	cut-like 1; CCAAT displacement	222.2758	34		330.5149		5.72E-01			Other
Q30201	H05ab3	hemochromatosis	205.1607	29		339,4607	278.446	7.26E-01			Other
P52895	116gh6	aldo-keto reductase family 1; me		26		339,7988		-3.56E+00			Other Other
Q9Y5N1	M24cd7	histamine receptor H3	526,5133	20		341,1301		-6.26E-01 -1.88E+00			Other
P43378	N23ab8	protein tyrosine phosphatase; no		44 40		368,4734 395,8948		-6.49E-01			Other
P35414	O11ab2	angiolensin receptor-like 1 FXYD domain-containing ion tran	620.7039	71		402.87			-1.61E+00		Other
Q14802 Q9Y266	K03cd1 F16cd6	nuclear distribution gene C homo		44		408.3037		-8,68E-01		20	Other
P30043	O21ab3	biliverdin reductase B (flavin redu		56		417.1383	620.0433	-1.07E+00	-6.35E-01	20	Other
Q9NP73	L06gh3	uncharacterized hematopoietic si		44	16	421.5183	557.2055	-9.32E-01	-8.51E-01		Other
Q12972	D17ab8	protein phosphatase 1; regulator	251.1796	35		425.52		7.61E-01	5.13E-01		Other
Q9Y296	N21ef2	PTD009 protein	738.5086	42		436.6114			-8.01E-01		Other
Q02750	K02ef7	mitogen-activated protein kinase	810.206	50		441.4249			-6.72E-01		Other Other
P42226	G23cd2	signal transducer and activator o			16	450.4229 452.267			-7.92E-01 -1.99E+00		Other
P31947	M07ef6	stratifin	2941.197	74	37	466.9229			-2.35E+00		Other
P46597	J12ab2	acetylserotonin O-methyltransfer			37 42	504.4269			-1.52E+00		Other
014745	J24cd4	solute carrier family 9 (sodium/h) G protein-coupled receptor 27	1464.475		31	595.0192			-1.94E+00		Other
Q9NS67	D24gh1	programmed cell death 6 interact			86	605.9272			-1.68E-01		Other
Q9UKL5		liver-specific bHLH-Zip transcrip		11		606,7901		-4.44E-01		20	Other
Q9NZZ1	P18ef2	hypothetical protein HSPC138	397,5554	4	50	618,4035	488,5944	6.37E-01	1.78E-01	20	Other
Q9Y3Q3		integral type I protein	526.2023	3	55	631,4284	504.2124		-5,68E-01		Other
Q9NSN3		similar to aspartate beta hydroxy			73	667.4052			-3.79E-01		Other
Q9Y5Z0	B12cd7	beta-site APP-cleaving enzyme:			60	680,3343			-6.14E-01		Other
	C21gh7	H2B histone family; member B	511.589		55	733,4035			7.41E-01 7.42E-01		Other Other
014713	M11cd5	integrin cytoplasmic domain-ass			46 60	737.9643 760.4217			-2.34E+00		Other
P46095	P11ab4	G protein-coupled receptor 6	2336.841 1761.627		53	779,7338			-1.43E+00		Other
Q9BV10		hypothetical protein MGC3136 calcium binding protein P22	1713.549	14		793,2383			-2.59E-01		Other
Q99653	O08cd7	polymerase (RNA) II (DNA direc			92	821.9856				20	Other
P41584 P11940	P16gh1 E06ab8	poly(A) binding protein; cytoplas	1738.585		69	827.8343			-1.38E+00	20	Other
Q00341	G21ab6	high density lipoprotein binding	545.5844	4	44	866,5328	618.5713	6.67E-01	-2.99E-01		Other
Q16795	K08ab7	NADH dehydrogenase (ubiquino		11	90	873.0478			-4.15E-01		Other .
Q13082	E05ab7	microtubule-associated protein			72	873.8683			-1,37E+00		Other
P36405	D16ab2	ADP-ribosylation factor-like 3	508.3031		47	939.7284					Other Other
Q9NXH		hypothetical protein FLJ20254	515.6731		67	959.9781			-1.44E-01 9,59E-01		Other
Q9Y323		hypothetical protein HSPC177	838.9663 Ii 560.9674		30 18	976.2242 1003.945					Other
P48059	112ab7	LIM and senescent cell antigen- milochondrial ribosomal protein	1 579 0093		36	1024.359					Other
Q9P0M9		spectrin; alpha; non-erythrocytic			86	1070.984			-1.35E+00	2	0 Other
Q13813 Q9Y379		CGI-69 protein	2556.713		00	1174.107			-4.26E-01	2	0 Other
Q13247			ic 718.1614	12	10	1194.951	1041.827				0 Other
P14416		dopamine receptor D2	814,5351		45	1299.03			-1.28E-01		0 Other
Q06323		proteasome (prosome; macropa			50	1386,719		-1.05E-01			0 Other 0 Other
Q9NR3		SAR1 protein	752.6343		86	1386.849			2.36E-01 -5.48E+00		0 Other
P05451	C07cd3	regenerating Islet-derived 1 alpha	# /3044.12		90 90	1394.425			-1.51E+00		0 Other
P04075		aldolase A; fructose-bisphospha ATPase; H+ transporting; lysos			80						0 Other
Q15904		NAD(P)H dehydrogenase; quin			303	1486.4	5 775.5735		-8.25E-01		0 Other
P15559 O00303		eukaryotic translation initiation			36	1556.44	8 1145,377	7 3.23E-01	1 -9.68E-01		0 Other
P56705					90	1635.80	5 2271.364	-1.13E+00	-1,18E+00		0 Other
P39210		MpV17 transgene; murine home	ol 990.336		394	1723.0	6 1202.587	7.99E-0	1 -1,47E-01		0 Other
Q9UKL	9 N03ef4	aldo-keto reductase family 1; m	e 16311.31		340		2 7016.79	3.10E+00	-2.52E+00		0 Other
043516	E12cd3	Wiskelt-Aldrich syndrome prote	ir 1162.511		190				3.51E-02		0 Other
Q9Y47		stem cell growth factor; lympho	cy 1346.693	11	110	2049.54	2 1503.55	5 6.06E-0	1 -2.73E-01		0 Other
P29312			1596.032		750 750	2331.85	9 4074.00	6 86EA	0 -1.40E-01 1 1.32E-01		0 Other
Q9Y5R					380			8 -1.15E+0	0 -1.07E+00		0 Other
P17676 O95362			atı 10407.52			2912.15			0 -1.96E+00		0 Other
O60417			1 6955.642		590	3097.57	9 5248.93	9 -1.17E+0	0 -2.89E-01	2	0 Other
P19623			1644.266		020	3417.97	8 2027.64	3 1,06E+0	0 -6.88E-01		0 Other
Q9P03		FXYD domain-containing lon to	ar 2410.08	2	120	4486.02	9 3004.62	7 8.96E-0	1 -1.87E-01		0 Other
Q9BUF		tubulin bela-5	2252.623	1	580	4730.75	5 2855.74	4 1.07E+0	0 -5.08E-01		0 Other
P0099	C23cd2	serine protease inhibitor; Kazal	67989.21	11	300	5416.17	5 28226.5	8 -3.65E+0	0 -2.59E+00		0 Other 0 Other
P11142		heat shock 70kD protein 8	11154.99	7	84C	5433.84	3 8144.60	1 -1.U4E+0 5 .2 ccc ^	0 -5.08E-01		to Other
P10620		microsomal glutathione S-trans	is 6/05,736					J +2.00E+U 4 -4.70E+∩	1 -1.47E+00 0 -1.91E+00		O Other
Q1334			4356.01	5	4 IC	5660.75			1 -1.08E+00		0 Other
P1588				2	tur	714A N7	5 4295.78	4 1.19E+0	0 -2.72E-01		20 Other
Q9HCI			oil 17951.73	. 11	800	8567.73	6 12775.9	3 -1.07E+0	0 -6.04E-01		20 Other
Q0494 Q9UPI			1 4625.515	8	280	8998.78	7 7300.38	7 9.60E-0	1 8.39E-01		20 Olher
P1398			ld 17383.17	19	200	10453.4	3 15673.3	8 -7.34E-0	1 1.42E-01		20 Other
P1595			74972.46	7	610	11417	.5 31332.7	4 -2.72E+0	10 -3.30E+00		20 Other
P0468		l tubulin; alpha; ubiquitous	15943.82	7	260	15413.4	4 12871.6	7 -4.88E-0	2 -1.14E+00		20 Other
P0468			15832.93	7	590	24085.3	35 15834.4	9 6.05E-0	1 -1.06E+00	:	20 Other

Q9U109	L02gh4	13kDa differentiation-associated	1754,698	1990	2889.148	2210.196 7.19E-	1.79E-01	2 Unahaanad I Kab
Q9Y6H1		16.7Kd protein	5687.333	4050			02 -4.90E-01	3 Unchanged High
Q9P0U1		6.2 kd protein	2323,402	1990			01 -2.21E-01	
P80095	N10ab7	6-pyruvoyl-lelrahydropterin synth		2620		2471.592 -1.04E+		
Q92747		actin related protein 2/3 complex		3990		3972.89 -6.92E-		
015144	F08ab2	actin related protein 2/3 complex		21400		17538.09 3.00E-		
O15145		actin related protein 2/3 complex		3190		2683.638 -7.73E-		
P02570	A01cd7	actin; bela	66304.81	45400		50171.41 -7.74E-		3 Unchanged High
P02570	H12cd7	actin; beta	64949.74	42400		50643.44 -5.42E-		
P02570	P12cd7	actin; beta	21347.36	12200		16331.85 -4.70E-		3 Unchanged High
P02571	A06ab6	actin; gamma 1	69165.16	59100		64643.45 -7.47E-		3 Unchanged High
P12814	E23ab2	actinin; alpha 1	2400.607	2160			1 -1.50E-01	3 Unchanged High
043707	G05ab2	actinin; alpha 4	6999.13B	3810		4769.196 -1.00E+		3 Unchanged High
P53999	C05cd7	activated RNA potymerase It trans		4320			1 -5.13E-02	
P18848	103ef5	activating transcription factor 4 (t	7160.888	5420			02 -4.02E-01	
P53680	A07ef1	adaptor-related protein complex:		2530		2963.423 8.63E-4		3 Unchanged High
P29274	A18ef5	adenosine A2a receptor	1990.124	2960	2004.596	2318.367 1.05E-		3 Unchanged High 3 Unchanged High
P54819	A04ab2	adenylate kinase 2	2997.607	2520		2788.3 -7.20E-		
Q01518	O18cd6	adenylyl cyclase-associated prot		11000	7171.205			
P32889	C03ef6	ADP-ribosylation factor 1	6040.796	7220	7907.02	7056.851 3.88E-0		
P18085	P21ab2	ADP-ribosylation factor 4	4777.904	8880	8342.165	7331.81 8.04E-		
P26437	B02ab2	ADP-ribosylation factor 5	3739.477	4430		4052.262 9.17E-0		
Q9BZX4	L08gh6	AKAP-binding sperm protein ropp		2780			2.44E-01	3 Unchanged High 3 Unchanged High
P11766	K01ab2	alcohol dehydrogenase 5 (class I		1990	2854.318		1 -1.59E-01	3 Unchanged High
P05067	N03ef6	amyfold beta (A4) precursor prote	2846.85	4180	3918.595	3649,558 4,61E-0		3 Unchanged High
Q06481	K22ab2	amyloid bela (A4) precursor-like		3230	2290.191	2546.598 1.13E-0		
P07355	112ab2	annexin A2	74767.39	112000	97217.17	94734.13 3.79E-0		3 Unchanged High 3 Unchanged High
P08758	118ab2	annexin A5	4487.905	4270			1 -7.32E-02	
P55064	O08ab2	aquaporin 5	10851.57	14200	11650,24	12248,87 1.02E-0		
Q13520	O10ab2	aquaporin 6; kidney specific	2404.748	2350			02 -3.25E-02	3 Unchanged High 3 Unchanged High
P32391	D19ab2	ARP3 actin-related protein 3 hon		4280	3941.05	4171.013 -1.23E-0		3 Unchanged High
O43776	E18ab7	asparaginyl-tRNA synthetase	2570.968	3010		2811,634 1.51E-0		3 Unchanged High
P24539	N22ab2	ATP synthase; H+ transporting; r		3800	3228.203		1 3.86E-01	3 Unchanged High
P48201	P04ab2	ATP synthase; H+ transporting; r	5694.182	4370	4857.955	4974.135 -2.28E-0		3 Unchanged High
P05496	N24ab2	ATP synthase; H+ transporting; r		1760	2400.575		1 -3.06E-01	3 Unchanged High
Q06055	P02ab2	ATP synthase; H+ transporting; r		3530	7732.552		1 -7.12E-01	3 Unchanged High
O75947	P14ab2	ATP synthase; H+ transporting; r	2668,028	2520	1881.974		1 -8.37E-02	3 Unchanged High
P56385	P06ab2	ATP synthase; H+ transporting; r		2090		2462,524 -3.60E-0		3 Unchanged High
P56134	P10ab2	ATP synthase; H+ transporting; r		17200	14181.18	14327.12 2.91E-0		3 Unchanged High
P18859	P08ab2	ATP synthase; H+ transporting; r		4840		4633,559 4.16E-0		3 Unchanged High
075964	P16ab2	ATP synthase; H+ transporting; r	2021.591	2630	2009.026	2219,11 -8.996-0		3 Unchanged High
P25705	N14ab2	ATP synthase; H+ transporting; r	3657.512	2410	2967.207	3009,915 -3,02E-0		3 Unchanged High
P06576	N16ab2	ATP synthase; H+ transporting; r	4072.978	3280	2557.962	3305,047 -6.71E-0		3 Unchanged High
P36542	N18ab2	ATP synthase; H+ transporting; r	2479.5	2030	2436.284	2313,989 -2.54E-0		3 Unchanged High
P48047	P18ab2	ATP synthase; H+ transporting; r	4222.314	4100	3541.047	3955.838 -2.54E-0		3 Unchanged High
Q01814	P01el5	ATPase; Ca++ transporting; plas		3180	3242.707	3193.535 3.84E-0		3 Unchanged High
Q99437	A11ab3	ATPase; H+ transporting; lysosor		3870	2087.291	2817.803 -2.58E-0	1 6.33E-01	3 Unchanged High
P36543	A09ab3	ATPase; H+ transporting; lysosor		2920	1659.749	2145.908 -1.60E-0	1 6.57E-01	3 Unchanged High
075348	A15ab3	ATPase; H+ transporting; lysosor	2734.484	5660	2721.878	3704.42 -6.67E-0	3 1.05E+00	3 Unchanged High
Q16864	A21ab3	ATPase; H+ transporting; lysoson	6452.29	15900	7960.234	10101.59 3.03E-0	1 1.30E+00	3 Unchanged High
P27449	A05ab3	ATPase; H+ transporting; lysosor	17134.14	24300	13120.57	18185.04 -3.85E-0	1 5.04E-01	3 Unchanged High
000244	L14ab2	ATX1 antioxidant protein 1 homo	2833.957	3500	3977.547	3435.675 4.89E-0	1 3.03E-01	3 Unchanged High
014503	M19ab3	basic helix-loop-helix domain cor		4280	3254.631	4169,761 -6,13E-0		3 Unchanged High
060238	L11ab5	BCL2/adenovirus E1B 19kD Inter	1473.428	2420	3224.141	2371.407 1.13E+0		3 Unchanged High
P30536 P01884	L21ab3 C23ab3	benzodiazapine receptor (periph		2890	2326.211	2405.625 2.19E-0		3 Unchanged High
P22004	113ab5	bela-2-microglobulin	64872	105000	36055,95	58478.91 -8.47E-0		3 Unchanged High
095415	B02ef3	bone morphogenetic protein 6	5036,985	4710	3780.005	4509.317 -4.14E-0		3 Unchanged High
075531	K11ab3	brain protein 13 Breakpoint cluster region protein	7498.734	11200	5348,84	8032,295 -4.87E-0		3 Unchanged High
P02593	A10ab6	calmodulin 2 (phosphorylase kin:	2541.789		2924.826	2736.992 2.03E-0		3 Unchanged High
P07384	O24ef1	calpain 1; (mu/l) large subunit	1573.77	4130	4742.485	3482.215 1.59E+0		3 Unchanged High
P27797	P05ab5	calreticulin	3429,132 10259,42	3420 9700	2390.011 15437.35	3078.324 -5.21E-0 11797.33 5.89E-0		3 Unchanged High
	N06gh8	CaM-KII inhibitory protein	20298.71	28200				3 Unchanged High
P47756	N14ab4	capping protein (actin filament) n	4348.079		21760,79 5015.377	23425.99 1.00E-0 4604.9 2.06E-0		3 Unchanged High
P15086	H13ab5	carboxypeptidase B1 (tissue)	9766.062	27.1			1 3.38E-02	3 Unchanged High
P52952	D22ab5	cardiac-specific homeo box	2859.197	1740		3273,378 -8.50E+0		3 Unchanged High
P13862	M21ef1	casein kinase 2; beta polypeptide		3840	2282.546	2292.94 -3.25E-0		3 Unchanged High
014675	102ef6	CASP8 and FADD-like apoptosis		2500			6.25E-02	3 Unchanged High
P35221	C17ef7	catenin (cadherin-associated pro		4230		2092.229 -1.19E+0 3782.772 -2.08E-0		3 Unchanged High
P07858	E01ab5	cathepsin B	2480,797	3860				3 Unchanged High
P48509	K16ab4	CD151 antigen	7267.56		8768.731	3607,853 8.54E-0 9901,71 2,71E-0	6.37E-01 9.11E-01	3 Unchanged High
P25063	G22cdB	CD24 antigen (small cell lung car				15932.79 -1.36E+0	0.11E-01	3 Unchanged High
P16070	O21ef1	CD44 antigen (homing function a				10533.43 -1.49E+0		3 Unchanged High
P08962	O18ab4	CD63 antigen (melanoma 1 antig		11600	10016,29		2 3.04E-01	3 Unchanged High
014519	K04ef5	CDK2-associated protein 1	4231.191	4180	4763.013		1 -1.90E-02	3 Unchanged High 3 Unchanged High
P25763	A12ab6	cell division cycle 42 (GTP bindir				3426.736 -6.86E-0		3 Unchanged High
P49368	N17cd2	chaperonin containing TCP1; sut				2484.939 -8.46E-0		3 Unchanged High
P48643	H10ef4	chaperonin containing TCP1; sut		2330	1561.625	2042.12 -5.20E-0		3 Unchanged High
000299	M19ab6	chloride intracellular channel 1	6811.55	7240		6591,299 -2,50E-0		3 Unchanged High
Q9Y2Q7	N02cd8	chromosome 11 open reading fre				4610,942 5.41E-0		3 Unchanged High
P09496	F15ab5	clathrin; light polypeptide (Lca)	5758.632			6957,341 2.26E-0		3 Unchanged High
095832	L04gh1	claudin 1	3046.376			2138.383 -1.73E+0		3 Unchanged High
095471	A13ef1		2888.819	3350		2587.177 -9.25E-0		3 Unchanged High
P23528	M20ab5	cofilin 1 (non-muscle)	16021.95	18700		16255.64 -1.88E-0		3 Unchanged High
P16989	L08ef6	cold shock domain prolein A	2942.723			2515,212 -5.08E-0		3 Unchanged High
P38936	021ef5	cyclin-dependent kinase inhibitor	3643.18		3437.076	3777.57 -8.40E-0		3 Unchanged High

Q9U114	B14ef1	cyclin-E binding protein 1	26180.8	29800	39655.02	31877.4	5.99E-01	1.87E-01	3 Unchanged High
P04080	G16ab3	cystatin B (stefin B)	4312.168	5420	2370.13	4034,272	-8.63E-01	3.30E-01	3 Unchanged High
P21291	J01ab5		2478,366	1710	2135,86		-2.15E-01		3 Unchanged High
000622	J15ef7		2070,203	1620	2978.542	2224.339		-3.50E-01	3 Unchanged High
P13073	L19ab5	cytochrome c oxidase subunit IV	6219,628	8550	6571.848	7115.048	7.95E-02		3 Unchanged High
P12074	K13gh1	cytochrome c oxidase subunit VII	3253.24	5800	4244.557	4434,13	3,84E-01	8.35E-01	3 Unchanged High
Q02221	E17ab6	cytochrome c oxidase subunit VII	2968.24	2820	4604.241	3464.904	6.33E-01	-7.28E-02	3 Unchanged High
P14854	A20ab6	cytochrome c oxidase subunit VII	1991.91	2660	2380.795	2344.331	2.57E-01	4.17E-01	3 Unchanged High
P09669	E12ab6	cytochrome c oxidase subunit VIv	2580,598	2090	1927.002	2197.58	-4.21E-01	-3.08E-01	3 Unchanged High
P14406	G02ab5	cytochrome c oxidase subunit VII		4500	4245,478	4530,346	-1,91E-01	-1.07E-01	3 Unchanged High
O14548	G23cd5	cytochrome c oxidase subunit VII	2182,954	2450	3040,586	2556.481	4.78E-01	1.64E-01	3 Unchanged High
P24311	K17ab5	cytochrome c oxidase subunit VII	3224,558	5150	3801.752	4057.995	2.38E-01	6.75E-01	3 Unchanged High
P15954	P04ab5	cytochrome c oxidase subunit VII	12335.74	12200	9883,688	11482.48	-3.20E-01	-1.27E-02	3 Unchanged High
P10176	B10ab5	cytochrome c oxidase subunit VII	3019,162	1900	3329.281	2750.974	1,41E-01	-6.65E-01	3 Unchanged High
O43293	K19ab4	death-associated protein kinase	2535.403	1850	2088.808	2157.783	-2.80E-01	-4.55E-01	3 Unchanged High
P46966	G14ef6	defender against cell death 1	7921.682	11200	9546.896	9563.241	2.69E-01	5,02E-01	3 Unchanged High
Q09753	G01ab6	defensin; bela 1	4071.982	7630	2153.638	4619.611	-9.19E-01	9.07E-01	3 Unchanged High
P18282	F23ab2	destrin (actin depolymerizing fact	7729.126	7570	6083,408	7126.01	-3.45E-01	-3.09E-02	3 Unchanged High
P07108	H04gh1	dlazepam binding inhibitor (GAB.	2541.011	1730	2995.768	2422.578	2.38E-01		3 Unchanged High
P31689	D18ab7	DnaJ (Hsp40) homolog; subfamil	2491.051	2930	1789.662	2402.398	-4.77E-01	2.32E-01	3 Unchanged High
Q9Y5T4	l10cd8	DNAJ domain-containing	2324.933	2520	2397.896	2414.457	4.46E-02	1.17E-01	3 Unchanged High
Q9Y463	G15cd5		8211.965	6870	5987,39	7023.599	-4.56E-01		3 Unchanged High
Q9NP97	C02ef8	dynein light chain 2A	4080.949	4160	5365.928	4534.283	3.95E-01	2.63E-02	3 Unchanged High
Q15701	L08cd3	dynein; cytoplasmic; light polyper		5800	4332.161		-5.25E-02		3 Unchanged High
P78545	O16ab4	E74-like factor 3 (ets domain trar	3781.196	2780	1455.944		-1.38E+00		3 Unchanged High
O60869	A03cd4	endothelial differentiation-related		5680	4090.527		-4.31E-01		3 Unchanged High
Q14259	L05ab4	enhancer of rudimentary homolog		3950	5175.131	4241.036	5.26E-01	1.38E-01	3 Unchanged High
P06733	C16ab4	enolase 1; (alpha)	12539.97	7460	9955.024	9983.774			3 Unchanged High
P22712	D11gh6	enolase 1; (alpha)	6900.857	4290	5016,695	5403.444			3 Unchanged High
P04720	007ab6	eukaryotic translation elongation eukaryotic translation elongation	36222.34	23200	23485.94		-6,25E-01 7,58E-01		3 Unchanged High
P26641 P13639	A10ab4		6863,565	6540	11607.61 3360.073	8337,901			3 Unchanged High
	115ab5 L24cd3	eukaryotic translation elongation eukaryotic translation initiation fa	4023.855	2310 1930	2391.869		-2.60E-01 -2.63E-01		 Unchanged High Unchanged High
015372	L22cd3		2870.027	2670	3333,728	3005.577		-1.70E-01	3 Unchanged High
O75821 Q64252	L11ab6	eukaryotic translation initiation fa eukaryotic translation initiation fa		1540	2127.874	2141.68			3 Unchanged High
015371	L18cd3		4273.919	4030	3558,838	3954.374			3 Unchanged High
P04765	A22ab4	eukaryotic translation initiation fa		7170	6363.169	7083.99			3 Unchanged High
Q14240	H13ab4	eukaryotic translation initiation fa		2190	2519.322	2267.015			3 Unchanged High
P10159	A18ab5	eukaryotic translation initiation fa		5240	5651.752		-6.75E-01		3 Unchanged High
Q9NQT4	H05gh4	exosome component Rrp46	2098.514	2590	2304.48	2331.056	1.35E-01		3 Unchanged High
P02794	C12ab6	femilin; heavy polypeptide 1	35912.81	48700	42075.22	42243.59		4.41E-01	3 Unchanged High
P02792	M10ab3	ferritin; light polypeptide	44225.86	29600	17849.2		-1.31E+00		3 Unchanged High
Q05472	B02ab4	Finkel-Blskls-Reilly murine sarco	10322.94	10400	9704.642	10145.46	-8.91E-02	1.19E-02	3 Unchanged High
P20071	F08ab3	FK506 binding protein 1A (12kD)	2741,389	1870	1414,969	2009.291	-9.54E-01	-5.51E-01	3 Unchanged High
P50395	J02ab4	GDP dissociation Inhibitor 2	2287.713	1580	2291.27	2051.41	2.24E-03	-5.38E-01	3 Unchanged High
P48507	E12ab5	glutamate-cysteine ligase; modifi	154.1773	76.2	5937.375	2055.914	5.27E+00	-1.02E+00	3 Unchanged High
P36969	D14ab5	glutathione peroxidase 4 (phospt	3397.2	4860	3108.036	3787.13	-1.28E-01	5.15E-01	3 Unchanged High
P09211	L10ab3	glutathione S-transferase pi	6392.334	8670	4067.789	6378.198	-6.52E-01	4,40E-01	3 Unchanged High
P04408	A01cd8	glyceraldehyde-3-phosphate deh	28809,37	38300	44655.67	37260.67	6.32E-01	4.11E-01	3 Unchanged High
P04406	H12cd8	glyceraldehyde-3-phosphate deh	37562.25	60200	57135.41	51621.55			3 Unchanged High
P04406	P12cd8	glyceraldehyde-3-phosphate deh	12053.5	14900	19061.62	15342.7	6.61E-01	3.07E-01	3 Unchanged High
P43304	H01ab3	glycerol-3-phosphate dehydroger		2340	3276.264	3023.392	_		3 Unchanged High
P41250	F02ef7	glycyl-IRNA synthetase	4333.439	3630	6248,604	4736.632		-2.56E-01	3 Unchanged High
P04895	K17ef5	GNAS complex locus	7755.858	9290	9107,558	8718.93		2.61E-01	3 Unchanged High
P24522	B03ef6	growth arrest and DNA-damage-i		6130	3396.736		2.83E-01		3 Unchanged High
Q92847	M24ab5	growth hormone secretagogue re		2100	2597.123		-1.23E-01		3 Unchanged High
P04901 P2538B	B01ef1	guanine nucleotide binding prote guanine nucleotide binding prote		7610 15300	5853,112 19055,7	6398.85	2.96E-02 -2.35E-01		3 Unchanged High
	E20cd6 G08ab6	H3 histone; family 3B (H3.3B)		2770	2042.864		-2.55E-01		3 Unchanged High 3 Unchanged High
P06351 P04792	A16ef5	heat shock 27kD protein 1	2448.264 41952.14	34500	31970.86	36135.19		1.80E-01	3 Unchanged High 3 Unchanged High
P11021	H01ef1	heat shock 70kD protein 5 (glucc		1600	4101.742	2976.537		-1.01E+00	3 Unchanged High
P38646	O19ab7	heat shock 70kD protein 9B (mor		2180	2220,509		-3.61E-01		3 Unchanged High
Q9UK76	J14ef1	hematological and neurological e		35400	41099.58	34286,74		4.27E-01	3 Unchanged High
P09651	F12ab7	heterogeneous nuclear ribonucle		1650	2524.082		1.05E-03		3 Unchanged High
Q9Y4J5	123ef1	helerogeneous nuclear ribonucle		2370	3770.457		-2.11E-02		3 Unchanged High
P09429	F10ab7	high-mobility group (nonhistone (1660	3502.047		4.67E-01		3 Unchanged High
P49773	M03el6	histidine triad nucleotide binding	6025,978	7790	7379.267	7064,436			3 Unchanged High
Q92769	N05ab4	histone deacelylase 2	2454.324	2220	1789,508		-4.56E-01		3 Unchanged High
P17693	F08ab7	HLA-G histocompatibility antigen	2048,009	2780	1844.177	2225.106		4,42E-01	3 Unchanged High
P17483	C15gh2	homeo box B4	2892,488	3490	2902.69	3095.004		2.71E-01	3 Unchanged High
Q9UL99	C01cd8	hyaluronoglucosaminidase 4	3623.461	2050	2189.656	2621.286	-7.27E-01	-8.21E-01	3 Unchanged High
Q9GZT3	L18gh7	hypothetical protein DC50	3343,922	3750	4298,676	3797.943			3 Unchanged High
Q9NWY5		hypothetical protein FLJ20533	3104.471	5460	6184.227	4915.419	9.94E-01		3 Unchanged High
Q9U130	P08ef2	hypothetical protein HSPC152	3000.217	4060	4396.862	3819.313	5.51E-01	4,37E-01	3 Unchanged High
Q9BQB6	P08gh5	hypothetical protein IMAGE3455:	3599.509	5410	8687.307	5897,337	1.27E+00		3 Unchanged High
Q9BWJ5	P18gh7	hypothetical protein MGC3133	2074.073	2520	2719.671	2436.767			3 Unchanged High
Q9Y683	N11ef2	hypothetical protein MGC8721	2425.475	2730	2939.253		2.77E-01	1.73E-01	3 Unchanged High
Q16665	P01ef6	hypoxia-inducible factor 1; alpha	3544.753	2960	1526.935	2675.668	-1.22E+00	-2.62E-01	3 Unchanged High
O75353	G04ef6	immediate early response 3	4564.408	4690	3129.721	4129.098	-5.44E-01	4.01E-02	3 Unchanged High
Q16270	J15ab6	insulin-like growth factor binding	6185.622	21800	18047.34	15334.47	1.54E+00	1.82E+00	3 Unchanged High
Q9Y287	A03gh2	integral membrane protein 2B	6908.997	10500	10014.47				3 Unchanged High
P56537	D16ab6	integrin beta 4 binding protein	2594.316	3270	1568.747		-7.26E-01	3,32E-01	3 Unchanged High
P26006	D08ab6	integrin; alpha 3 (antigen CD49C		2740	1339.712		-9.95E-01		3 Unchanged High
P05556	G01ef7	integrin; beta 1 (fibronectin recep		5390	6511.579		-5.05E-02	-3.22E-01	3 Unchanged High
Q01628	C09gh7	interferon induced transmembrar		14800	8827.997	12669.24	-7.09E-01	3.23E-02	3 Unchanged High
Q12905	C10ab7	Interleukin enhancer binding fact	3045.484	2420	2596,915	2687.538	-2.30E-01	-3.32E-01	3 Unchanged High

Q13907	A24ab7	isopentenyl-diphosphate delta is:		1660	2475,365	2362.933	-2.57E-01	-8.38E-01	3 Unchanged High
P33947	K03cd7	KDEL (Lys-Asp-Glu-Leu) endoph	1667.246	1250	3924.993	2281.692	1.24E+00	-4.12E-01	3 Unchanged High
P00338	D15ab7	lactate dehydrogenase A	44081,29	34500					3 Unchanged High
P08865	G21e/7	taminin receptor 1 (67kD; ribosor		6070	9921.409	8180.607	2.15E-01	-4.93E-01	3 Unchanged High
Q08380	D17ab7	lectin; galactoside-binding; solub		2850	3014.02	2719.794	3.94E-01		3 Unchanged High
014949	F17ef3	low molecular mass ublquinone-t		3550	3637,463		3.06E-01	2.71E-01	3 Unchanged High
Q15012	F20ef4	hysosomal-associated protein trai		4690	5453,634	4319.567	9.54E-01	7.36E-01	3 Unchanged High
P14174	J05ef7	macrophage migration inhibitory		6700	6763.718	6337.062	2.87E-01	2.74E-01	3 Unchanged High
P49006	M05efB	macrophage myristoylated alanir		10400	8986.315		6.17E-01	8.22E-01	3 Unchanged High
P30463	G10ab6	major histocompatibility complex;		4890	2255.929		-2.60E-01	8.55E-01	3 Unchanged High
P10321	A01cd2	major histocompatibility complex;		17000	10196.88	12044.89	1.88E-01	9.25E-01	3 Unchanged High
P10321	H12cd2	major histocompatibility complex;		21600	11111.54	13298.19	6.28E-01		3 Unchanged High
P10321 P13747	P12cd2	major histocompatibility complex;		16800	11725.68		-1.62E-01	3.53E-01 1.33E-01	3 Unchanged High
Q95HC0	M24ab7 P02gh6	major histocompatibility complex; major histocompatibility complex;		2790 4140	2053,351 3295,115		-3.07E-01 -6.21E-01		3 Unchanged High 3 Unchanged High
P13640	D13gh6	metallothionein 1G	6180.376	4750	2940.683		-1.07E+00	-2.90E-01 -3.79E-01	3 Unchanged High
P80297	D17gh6	metallothionein 1X	4881.982	5310	5233.749	5142,998	1.00E-01	1.22E-01	3 Unchanged High
P02795	L04ab7	metallothionein 2A	6620,235	7330		7340.215	2.85E-01	1.48E-01	3 Unchanged High
014880	E04ab7	microsomal glutathione S-transfe		7940	4177.165		-2.66E-01	6.60E-01	3 Unchanged High
Q9NYZ2	M22ef2	mitochondrial solute carrier	2488.07	2650	2240.824		-1.51E-01	1,98E-01	3 Unchanged High
P26038	K05ab7	moesin	4260,51	4580	3801.298	4212,427	-1,65E-01	1.03E-01	3 Unchanged High
015329	O12ef1	mucin 6; gastric	5574.093	5950	6685.047	6068.269	2.62E-01	9.31E-02	3 Unchanged High
P16475	J02gh1	myosin; light polypeptide 6; alkal	48122.62	45200	48882.54	47407.02	2.26E-02	-8.99E-02	3 Unchanged High
P19105	N03cd6	myosin; light polypeptide; regulat	7920.873	8150	7228.401	7764.994	-1.32E-01	4.04E-02	3 Unchanged High
Q9UK23	D06ef1	N-acetylglucosamine-1-phosphoi	1884.596	2680	2511.293	2357.087	4.14E-01	5.05E-01	3 Unchanged High
015239	E22ab7	NADH dehydrogenase (ubiquino		5440	3719.503	4005.894	3.79E-01	9.27E-01	3 Unchanged High
000483	J02ab7	NADH dehydrogenase (ubiquinor		5540	9086.615	6613.046	8.01E-01	8.70E-02	3 Unchanged High
075438	J04ab8	NADH dehydrogenase (ubiquino		2170	2842.647	2212,932	8.04E-01	4.13E-01	3 Unchanged High
095168	J08ab8	NADH dehydrogenase (ubiquinor		3270	2253,854	2607.423		5.09E-01	3 Unchanged High
095298	J12ab8	NADH dehydrogenase (ubiquino)		2840	2888.281	2544.679	5.97E-01	5.71E-01	3 Unchanged High
043920	J18ab8	NADH dehydrogenase (ubiquinos		11500	8014.187	8594.08	3.61E-01	8.85E-01	3 Unchanged High
Q13765 Q15843	H09ab7 K13cd1	nascent-polypeptide-associated in neural precursor cell expressed:		3390 4710	3897.956 4139.56	3363.371 4098.747	4.77E-01 2.63E-01	2.76E-01	3 Unchanged High
Q9NX14	C01gh3	neuronal protein 17,3	3449.484 3476.812	3150	5077.834	3902.8		4.48E-01 -1.41E-01	3 Unchanged High 3 Unchanged High
Q9Y2R6	F18gh6	NICE-3 protein	1763.974	2250	2329.521	2113.511	4.01E-01	3.49E-01	3 Unchanged High
Q15668	H13cd6	Niemann-Pick disease; type C2	1516.592	3640	2828.255	2662.845	8.99E-01		3 Unchanged High
014597	J21ef4	non-functional folate binding prof		2170	3004.012			-4.27E-01	3 Unchanged High
P15531	E02ef5	non-metastatic cells 1; protein (N		6940	7836.897	7143.349	2.36E-01	6.07E-02	3 Unchanged High
P22392	L12cd2	non-metastatic cells 2; protein (N		5680	4836.596	5114.664	2.72E-03	2,35E-01	3 Unchanged High
000746	D04ef7	non-metastatic cells 4; protein ex		2270	4300,604	2927,223	9.60E-01	3.80E-02	3 Unchanged High
075534	B01cd4	NRAS-related gene	3478.657	3510	2737.425	3241.269	-3.46E-01	1.20E-02	3 Unchanged High
Q9H9A1	C24gh6	nuclear receptor co-repressor/HC	5951.715	3290	2990.083	4076,544	-9.93E-01	-8.56E-Q1	3 Unchanged High
Q9NPE3	K15ef4	nucleolar protein family A; memb		7130	3639,205			1.01E+00	3 Unchanged High
P54368	G10ef1	ornithine decarboxylase antizym		6880	7782,339		1.10E-01		3 Unchanged High
Q9UH52	L23cd8	over-expressed breast tumor pro-		2320	1884.264		-3.10E-01		3 Unchanged High
Q15070	A09cd1	oxidase (cytochrome c) assembly		2500	1436.463			2.08E-01	3 Unchanged High
Q9H230 O60356	G14gh5 M03cd8	p53-induced protein PIGPC1 p8 protein (candidate of metasta:	1823.554	3330 1940	1443,914			8.69E-01	3 Unchanged High
P78337	M22ab8	paired-like homeodomain transcr		13300	2740.792 14490.63	2150.631 14411,94	6,27E-01 -9.05E-02	1.25E-01	3 Unchanged High 3 Unchanged High
Q9C086	P20gh7	PAP-1 binding protein	2992.2	3610	5014.215	3872.969	7.45E-01	2.72E-01	3 Unchanged High
Q15165	D06ef6	paraoxonase 2	1325.972	3740	2122.119			1.49E+00	3 Unchanged High
P20962	N13ab8	parathymosin	2345,869	2330	3489.616			-8.88E-03	3 Unchanged High
P23284	117ab8	peptidylprolyl Isomerase B (cyclo		5860	8890.43		7.47E-01	1.45E-01	3 Unchanged High
Q06830	E18ab8	peroxiredoxin 1	2973.862	2450	2056.112		-5.32E-01		3 Unchanged High
P32119	G04ef7	peroxiredoxin 2	3141,718	3330	2921.471	3130.835	-1.05E-01	8.37E-02	3 Unchanged High
P30048	N09ab2	peroxiredoxin 3	2420.936	2010	1671.796	2035.218	-5.34E-01	-2.66E-01	3 Unchanged High
Q13162	N07ab2	peroxiredoxin 4	2744.707	2080	3645.718	2822.139	4.10E-01		3 Unchanged High
P30044	P15cd7	peroxiredoxin 5	2205,14	3820	1714.041	2581.24		7,94E-01	3 Unchanged High
Q13492	B13cd4	phosphatidylinositol binding clatt		2690	1861.508			3.03E-03	3 Unchanged High
P08237 P00558	P02ab7 P06ab7	phosphofructokinase; muscle	2790.708	1980	2829,743			-4.94E-01	3 Unchanged High
P18669	K14ab8	phosphoglycerate kinase 1 phosphoglycerate mutase 1 (brai	5186.346	5130	6341.68 2859.446	5552.736 2477.75		-1.57E-02	3 Unchanged High
Q14801	B01cd8	phosphoprotein enriched in astro		2730 17400	19729.36		6.30E-01	5.61E-01	3 Unchanged High
014832	A04cd1	phylanoyl-CoA hydroxylase (Ref:		7780	8341.917	7819.036	-2.39E-01 1.86E-01		3 Unchanged High
P53801	L01ef1	pituitary tumor-transforming 1 Int		5330					3 Unchanged High 3 Unchanged High
Q9HB21	E19gh5	pleckstrin homology domain-cont	2318.08	3070	2907.381	2766.359			0.14-1
Q15365	O09cd1	poly(rC) binding protein 1	4617,635			3635.431			3 Unchanged High
P52433	B17ab8	polymerase (RNA) II (DNA direct		3780		3198,285			3 Unchanged High
P52436	L16gh1	polymerase (RNA) II (DNA direct		2600	2709,495		3.26E-01		3 Unchanged High
Q03052	B19ab8	POU domain; class 3; transcriptie		2650	2251.677		-5.61E-01		3 Unchanged High
	K20gh5	PP1201 protein	7127.293	6650	2688.636		-1.41E+00		3 Unchanged High
P40425	G08ab8	pre-B-cell leukemia transcription		2650	3413.81	2958,976	2.77E-01	-9.05E-02	3 Unchanged High
Q99471	K06ab8	prefoldin 5	1827.777	2160		2275.038	6.34E-01	2.41E-01	3 Unchanged High
Q9UHZ2	E21ef8	PRO1073 protein	1872.208	3140	1690.101		-1.48E-01		3 Unchanged High
P07737	A21cd1	profilin 1	29650.38	48900	35394.8	37970.06	2,55E-01		3 Unchanged High
Q92740	E03gh1	prosaposin (variant Gaucher dist		5370	4268.459	4622.898	1.13E-02		3 Unchanged High
P07478	NO2cd2	protease; serine; 2 (trypsin 2)	77359,27		51/45.68	57240.67	-5.80E-01	-8,60E-01	3 Unchanged High
Q9UL46 P25786	N09ab8 L09ef7	proleasome (prosome; macropali		4550		3648.995			3 Unchanged High
P20618	J17ab8	proteasome (prosome; macropais proteasome (prosome; macropais		3590	2265,763	3082,003	-2.75E-03	1.80E-01	3 Unchanged High
P49720	J21ab8	proteasome (prosome; macropali		2100			-7.52E-03 5.06E-01		3 Unchanged High 3 Unchanged High
P28070	J23ab8	proteasome (prosome; macropair	2193 138	1760			2.04E-02		3 Unchanged High
Q99436	L03ab8	proteasome (prosome; macropair		3450			-4.44E-01	-2 785-01	3 Unchanged High
P08129	G01ef6	protein phosphalase 1; catalytic 1		2200		2339.797	-1.60F-01	-2.09F-01	3 Unchanged High
015355	D07ab8	protein phosphatase 1G (former)		4200	3717.85		-6.65E-01		3 Unchanged High
P30153	A06ef3	protein phosphatase 2 (formerly):		4100	2765.011		-4.81E-01		3 Unchanged High

Q15249	H13ef5	prothymosin; aipha (gene sequer	11382.44	14500	9974.587	11954.68			3 Unchanged High
Q9NQ11	G03gh7	putative ATPase	2774.724		2271.593		-2.89E-01		3 Unchanged High
P41567	N05cd5	putative translation initiation facts			8909.545 2667.605	7666.779	2.51E-01 -8.82E-01		3 Unchanged High 3 Unchanged High
Q15181	L18gh1	pyrophosphatase (inorganic)	4914.897 6305,222		9000.504		5.13E-01	3.54E-01	3 Unchanged High
P14786 P57735	M24ab8 G12gh4	pyruvate kinase; muscle RAB25; member RAS oncogene	3343.628	4890	1366.379	3198.699		5.47E-01	3 Unchanged High
P54725	E21cd1	RAD23 homolog A (S. cerevisiae	2533.36	1790	1753.113		-5,31E-01	-5.00E-01	3 Unchanged High
P54727	M10ef6	RAD23 homolog B (S. cerevisiae		5410	4983.21	4809.132		4,20E-01	3 Unchanged High
000538	P16cd5	RAS guanyl releasing protein 2 (-			1906.199		-3.44E-01	-6.90E-02 -1.43E-01	3 Unchanged High 3 Unchanged High
P06749	C21gh1	ras homolog gene family; membe	2136,708	1940 4490	3003,841 2930,236	2358.668 3943.029		2,91E-02	3 Unchanged High
P08134	B16ab2 P06gh8	ras homolog gene family; membe ras-like protein VTS58635	5451.542		9066,798	7033.479	7.34E-01		3 Unchanged High
P15154	CO1ef6	ras-related C3 botulinum toxin su		9720	10044.19		-1.11E-01		3 Unchanged High
Q15347	B20cd6	Ras-related GTP-binding protein	1728.963		2623.418			4.63E-01	3 Unchanged High
O95197	A06cd6	reticulon 3	3712.756	3420	3833.411	3654.06 3023.27	4.61E-02		3 Unchanged High 3 Unchanged High
P09455	F12ab8	retinol binding protein 1; cellular Rho GDP dissociation inhibitor (t	2211.12 3166.04	4480 3200	2383,274 3141,434		-1.13E-02	1.02E+00 1.52E-02	3 Unchanged High
P52565 P27635	K13ef6 N19cd2	ribosomal protein L10	13406.29	7100	11103.69		-2.72E-01		3 Unchanged High
P53025	G22cd1	ribosomal protein L10a	19374.73	15100	19939.26	18124.91		-3.63E-01	3 Unchanged High
P39026	D07cd1	ribosomal protein L11	14172.29	6760	10346.92				3 Unchanged High
P30050	D09cd1	ribosomal protein L12	38376.69	24500	38669,34	33860.13	1.10E-02 2.43E-01		3 Unchanged High3 Unchanged High
P40429	A01ef1	ribosomal protein L13a ribosomal protein L14	23800.5 18507.36	22200 14600	28159.37 16108.18	16402.14			3 Unchanged High
P50914 P39030	B18cd4 C13cd3	ribosomal protein L15	8407.532	8100	10809.12			-5.43E-02	3 Unchanged High
P18621	D19cd1	ribosomal protein L17	8779.141	8440	10004.78	9074.168		-5.71E-02	3 Unchanged High
Q07020	D13cd1	ribosomal protein L18	12168.43	10200	15619,17	12658.45		-2.56E-01	3 Unchanged High
Q02543	C15gh1	ribosomal protein L18a	24976.59	27200	28170.19	26796.74 5762.718	1.74E-01 -1.70E-01	1.25E-01	3 Unchanged High 3 Unchanged High
P14118	D15cd1	ribosomal protein L19 ribosomal protein L21	6226,589 11156,59	5530 11800	5535.643 16742.21	13222.79		7.72E-02	3 Unchanged High
P46778 P35268	A13cd3 D17cd1	ribosomal protein L22	16289.45	16200	18991.35	17168.94		-5.63E-03	3 Unchanged High
P23131	F11cd4	ribosomal protein L23	20972.93	20400	25729.93			-4.24E-02	3 Unchanged High
P29316	G09cd3	ribosomal protein L23a	6385.675	4840	5446.963		-2.29E-01		3 Unchanged High
P38663	D21cd1	ribosomal protein L24	7805,803	6180	5479.195	9469.252	-5.11E-01	-1.29E+00	3 Unchanged High 3 Unchanged High
P08526 P46776	F01cd1 F05cd1	ribosomal protein L27 ribosomal protein L27a	12874.22 16046.37	5280 17000	10256.69 20949.85		3.85E-01		3 Unchanged High
P46779	F07cd1	ribosomal protein L28	40540.59	32400	45810.52			-3.24E-01	3 Unchanged High
P47914	E23cd3	ribosomal protein L29	3346.473	1960	1947.028			-7.73E-01	3 Unchanged High
P39023	B21cd1	ribosomal protein L3	13192.5	10500	13962.55			-3.36E-01 -5.20E-01	3 Unchanged High 3 Unchanged High
P04645	F03cd1	ribosomal protein L30	17693.61 9749.468	12300 8980	21530.83 12591.65			-1.18E-01	3 Unchanged High
P12947 P02433	F09cd1 A15cd3	ribosomal protein L31 ribosomal protein L32	22774.44	24400	30742.77				3 Unchanged High
P49207	F11cd1	ribosomal protein L34	13251.1	11500	13193.66	-		-1.99E-01	3 Unchanged High
P42766	K16cd7	ribosomal protein L35	10454.27	10500	10655.65				3 Unchanged High
P18077	F13cd1	ribosomal protein L35a	14583.91	10700	8811.726			-4.50E-01	3 Unchanged High 3 Unchanged High
P09896	J06gh1	ribosomal protein L36a ribosomal protein L36a-like	8395.644 5167.775	8780 5750	12052.18 4693.401		5.22E-01 -1.39E-01		3 Unchanged High
P09896 P02403	A19cd3 F15cd1	ribosomal protein L37	31895.4	45000	41301.54				3 Unchanged High
P12751	F17cd1	ribosomal protein L37a	18652.48	17300	19955.82			-1.05E-01	3 Unchanged High
P23411	F19cd1	ribosomal protein L38	5149.82	5570	7560.21				3 Unchanged High 3 Unchanged High
P02404	A17cd3	ribosomal protein L39	14114.05 14367.47	21300 10400	22827.16 10448.85			5.96E-01 -4.72E-01	3 Unchanged High
P36578 P28751	B23cd1 L06gh1	ribosomal protein L4 ribosomal protein L41	35135.75	34300				-3.32E-02	3 Unchanged High
P46777	E21cd3	ribosomal protein L5	12002.66	8880	10351.69			-4.35E-01	3 Unchanged High
Q02878	D01cd1	ribosomal protein L6	15861.7	7300				-1.12E+00	3 Unchanged High
P18124	D03cd1	ribosomal protein L7	21230.67	15900 8240	19538.32 10586.53			-4,18E-01 -7.00E-01	3 Unchanged High 3 Unchanged High
P11518 P25120	G11cd3 D05cd1	ribosomal protein L7a ribosomal protein L8	13385.5 23580.43	18300				-3.67E-01	3 Unchanged High
P32969		ribosomal protein L9	28853.42	32200				1.56E-01	3 Unchanged High
P46783	H13cd1	ribosomal protein S10	7453.342	7660					3 Unchanged High
P04643	H15cd1	ribosomal protein S11	13429.85	15600					3 Unchanged High 3 Unchanged High
P25398		ribosomal protein S12 ribosomal protein S13	16170.73 13956.88	10600 12600				-6.11E-01 -1.48E-01	3 Unchanged High
Q02546 P11174		ribosomal protein S15	21322.73	19600				-1.24E-01	3 Unchanged High
P39027		ribosomal protein S15a	16787.1	15100				-1.53E-01	3 Unchanged High
P17008	J01cd1	ribosomal protein S16	26604.5	15700				-7.62E-01	3 Unchanged High
P08708		ribosomal protein S17	13979.25	7990 12600				1 -8.07E-01 1 6.84E-03	 Unchanged High Unchanged High
P25232 P17075		ribosomal protein S18 ribosomal protein S20	12558.18 21944.13	19900				1 -1.43E-01	3 Unchanged High
P35265		ribosomal protein S21	14046.7	9470				1 -5.69E-01	3 Unchanged High
P39028		ribosomal protein S23	12228.96	10900				1 -1.72E-01	3 Unchanged High
P16632		ribosomal protein S24	12913.37	13100				2 1.53E-02	
P25111		ribosomal protein \$25	6274.159 26138.26	5630 30900				1 -1.55E-01 1 2,42E-01	3 Unchanged High 3 Unchanged High
P02383				12800				1 -4,51E-01	3 Unchanged High
P42677 P14798			3284,551	3180				1 -4.77E-02	3 Unchanged High
P25112	J15cd1	ribosomal protein S28	30384,24	32000				2 7.66E-02	
P30054	J17cd1	ribosomat protein S29	18454.55		25539.2			1 1.79E-01	
P23396			5913.682	2830				2 -1.07E+00 1 -2.06E-01	
P49241 P12750			7154.088 19821.56	6200 16200				1 -2.06E-01 1 -2.87E-01	
P12750			2629.697	2330				2 -1.72E-01	3 Unchanged High
P46782			. 15421.08	11000	17355.5	4 14598.9	6 1.70E-0	1 -4.85E-01	3 Unchanged High
P10660		ribosomai protein S6	46408.18	26100				1 -8.30E-01	
P23821			18906.94 15941.2	16200 12300				1 -2.19E-01 1 -3.76E-01	
P09058 P46781			6215,562					1 -6.63E-01	
, 40,0		proton os							

P46781	H12cd5	ribosomal protein S9	8827.416	9710	15295.98	11278.36	7.93E-01	1.38E-01	3 Unchanged High
P46781	P12cd5	ribosomal protein \$9	9059,523	1790	5725,082	5525.883	-6.62E-01	-2.34E+00	3 Unchanged High
P05387	H01cd1	ribosomal protein; large P2	16511.14	12700	17775.72	15695.14		-3.87E-01	3 Unchanged High
P05388	F21cd1	ribosomal protein; large; P0	17127.41	12400	19746,88	16430.54		-4.64E-01	3 Unchanged High
P05386	F23cd1	ribosomal protein; large; P1	2228.298	1880	3134.529	2413.78		-2.46E-01	3 Unchanged High
Q9Y254 Q9NWJ8	C08ef3	ring-box 1	2597.578	4260	3365,165	3408.892 2666.66	3.74E-01		3 Unchanged High
P08206	A23gh3 D06cd1	roundabout homolog 4; magic roi \$100 calcium binding protein Atl	1969.602 38485.73	3940 49800	2093,279 31745,47		8.79E-02 -2.78E-01	3.72E-01	3 Unchanged High 3 Unchanged High
P31949	E09cd3	S100 calcium binding protein A1	8890,594	11100	5408,269		-7.17E-01	3.20E-01	3 Unchanged High
P06703	P17gh6	S100 calcium binding protein A6	3403.146	4070	2088,478		-7.04E-01	2.60E-01	3 Unchanged High
P55735	J04ef4	SEC13-like 1 (S. cerevislae)	1695.152	1990	2362.541	2016.062	4.79E-01	2.32E-01	3 Unchanged High
P38384	106ef3	Sec61 gamma	5653,955	7820	7254.423	6908.432	3.60E-01	4.67E-01	3 Unchanged High
Q9NZJ3	H06ef2	selenoprotein T	2923.813	2440	1325.491	2230.091	-1.14E+00	-2.60E-01	3 Unchanged High
015532	H16cd1	selenoprotein W: 1	2678.843	2970	3918.44		5.49E-01		3 Unchanged High
Q13501	J23cd4	sequestosome 1	3214.395	3170	1164.17		-1.47E+00		3 Unchanged High
P35237 P05121	L18ab8 P05ef7	serine (or cystelne) proteinase in		8180	2938,72		-5.17E-01		3 Unchanged High
Q00587	C14cd7	serine (or cysteine) proteinase in serum constituent protein	4454.56 2480.698	7280 2660	9179.863 2044.608		1.04E+00 -2.79E-01	7.09E-01 9.80E-02	3 Unchanged High 3 Unchanged High
Q9H299	P16gh7	SH3 domain binding glutamic aci	14473,6	17100	13798.02	15134.74		2.43E-01	3 Unchanged High
P37108	E21cd2	signal recognition particle 14kD (35600	24319.59	28082.56	-3.96E-04	5.49E-01	3 Unchanged High
P49458	E19cd2	signal recognition particle 9kD	1874.967	2640	3141.422	2551.928	7.45E-01	4.93E-01	3 Unchanged High
P43308	G15cd2	signal sequence receptor; beta (I	4209.147	4610	5876.451	4898.581	4.81E-01	1.31E-01	3 Unchanged High
P51571	O15gh1	signal sequence receptor; delta (2062.113	2330	2654.968	2349.29	3.65E-01	1.77E-01	3 Unchanged High
000422	M17cd6	sin3-associated polypeptide; 18k	2502.79	2610	2079.295		-2.67E-01	5.83E-02	3 Unchanged High
075918	J10cd5	small EDRK-rich factor 2	5927.595	7010	7926.059	6955.245	4.19E-01	2.42E-01	3 Unchanged High
P13500	C15cd3	small inducible cytokine A2 (mon		38100	8786.323		-9.59E-01		3 Unchanged High
Q15356 P14648	A09cd2 A13cd2	small nuclear ribonucleoprotein p small nuclear ribonucleoprotein p		2150	2819.855	2570.901		-3.52E-01	3 Unchanged High
P12236	G01ef1	solute carrier family 25 (mitochor		2800 4760	2362.597 6176.034	2021.429 6107.995	1.39E+00		3 Unchanged High
Q00325	K22ab8	solute carrier family 25 (mitochor		3750	3779.577	4200,626		-6.32E-01	3 Unchanged High 3 Unchanged High
P32745	NO1cd1	somatostatin receptor 3	4102.668	4170	4748.989	4340.964	2.11E-01	2.39E-02	3 Unchanged High
P30626	E13cd2	sorcin	1895.592	2790	2280.297	2323,149	2.67E-01	5,59E-01	3 Unchanged High
Q01826	F18ef6	special AT-rich sequence binding		41300	24225,61		-1.27E-01	6,42E-01	3 Unchanged High
P21673	D12cd1	spermidine/spermine N1-acetyltn	8349.882	16500	6013.659	10275.93	-4.74E-01	9.80E-01	3 Unchanged High
P34991	G05cd3	S-phase kinase-associated prote	3072.601	2990	2396.506	2820,643			3 Unchanged High
P23246	B09cd2	splicing factor proline/glutamine		2460	3348.423	2834.056			3 Unchanged High
P35716	C07cd2	SRY (sex determining region Y)-I		4860	2125.089		-1.24E+00		3 Unchanged High
P35713	E10ef4	SRY (sex determining region Y)-I		3100	2024.386		-3.48E-01	2,66E-01	3 Unchanged High
Q06945	C05cd2	SRY (sex determining region Y)-I		2210	1636.067		-4.91E-01		3 Unchanged High
P00441 P50502	G20ef7 118cd2	superoxide dismutase 1; soluble	4439.342	4930 2530	3239,195		-4.55E-01	1.51E-01	3 Unchanged High
Q9Y5Y6	P20gh1	suppression of tumorigenicity 13 suppression of tumorigenicity 14	2307.599 2358.28	2700	2358.117 1494.856	2400.085	3.12E-02 -6.58E-01	1.35E-01 1.93E-01	3 Unchanged High
P17600	J06cd2	synapsin I	1969.157	2220	2010.908	2067,417	3.03E-02	1.74E-01	3 Unchanged High 3 Unchanged High
043759	G11cd5	synaplogyrin 1	14516.6	10700	10060.61	11775.53		-4.33E-01	3 Unchanged High
Q12962	P19cd2	TAF10 RNA polymerase II; TATA		5080	3962,323	4992,528	-5.83E-01		3 Unchanged High
Q15763	B20cd2	t-complex-associated-testis-expn		2890	2885.017	2600.459	5.12E-01	5.16E-01	3 Unchanged High
P10599	B23cd4	thioredoxin	11525.16	11900	6440,37	9965.777	-8.40E-01	5.00E-02	3 Unchanged High
P13472	002ef7	lhymosin; beta 10	46982.91	74300	54263.48	58508.68	2.08E-01	6.61E-01	3 Unchanged High
P01253	L12gh1	thymosin; beta 4; X chromosome		26000	16231.51		-5.12E-01	1.66E-01	3 Unchanged High
Q01085 Q15370	C02cd2 J12cd2	TIA1 cytotoxic granule-associate		1770	2917,342	2312.331		-3.47E-01	3 Unchanged High
043680	011cd2	transcription elongation factor B : transcription factor 21	1789.295	3980 1790	3919.327 3044.118	3505,384 2209,014	5.83E-01 7.67E-01	6.05E-01	3 Unchanged High
P21980	016cd2	transglutaminase 2 (C polypeptid		3490	1857.596		-7.53E-01	3.49E-03 1.58E-01	3 Unchanged High 3 Unchanged High
P29401	N15cd1	transketolase (Wernicke-Korsakı		2040	2715,206	2671.504	-2.63E-01	-6.75E-01	3 Unchanged High
060739	O01cd6	translation factor suit homolog	2671.935	3400	3868,727	3315,099	5.34E-01	3.50E-01	3 Unchanged High
Q9NS69	102gh7	translocase of outer mitochondris		2450	3754.914	3062.809		-2.89E-01	3 Unchanged High
P00938	M24cd1	triosephosphate isomerase 1	5701.29	5820	8345,738	6622.243	5,50E-01	2.97E-02	3 Unchanged High
P07226	G10cd2	tropomyosin 4	2636,256	2300	3755.882	2897.509	5.11E-01	-1.97E-01	3 Unchanged High
P04687	F09cd4	tubulin; alpha 3	20220.03	14500	22303,88	18992.31	1.42E-01		3 Unchanged High
P04687	A01cd1	tubulin; alpha; ubiquitous	15258.81	7840	22478,65	15193.76		-9.60E-01	3 Unchanged High
Q13509	E04cd6	tubulin; beta; 4	2338.877	2990	1853,273		-3.36E-01	3.56E-01	3 Unchanged High
P04350 075347	F01gh6 O08cd2	tubulin; beta; 5 tubulin-specific chaperone a	7573,765 3000,065	4430 3900	3448.455 3275.629	5149.498 3390.719	-1.14E+00 1.27E-01	-7.75E-01 3.77E-01	3 Unchanged High
075509	L10ef3	tumor necrosis factor receptor su		4890	1881,231		-2.55E-01		3 Unchanged High 3 Unchanged High
P13693	G16cd2	tumor protein; translationally-con		60800		49731.11			3 Unchanged High
P14625	G24cd2	tumor rejection antigen (gp96) 1	2185.187	2060	4905.342				3 Unchanged High
P42655	K04cd4	tyrosine 3-monooxygenase/trypic		5730	5558.974		-2.44E-01		3 Unchanged High
Q04917	G06cd3	tyrosine 3-monooxygenase/trypto		8150		7111.932			3 Unchanged High
P29312	A01cd4	tyrosine 3-monooxygenase/trypto	5873.282	4670	2279.148	4275.506	-1.37E+00	-3.29E-01	3 Unchanged High
P29312	P12cd4	tyrosine 3-monooxygenase/trypto		1140	1520.2		-1.59E+00		3 Unchanged High
014957	G17cd7	ubiquinol-cytochrome c reductas		3990		3351.45			3 Unchanged High
P14793	D01cd4 D22gh1	ubiquitin A-52 residue ribosomal			3258,439				3 Unchanged High
P02248 Q76069	B12cd4	ubiquitin B	2753.942		4152.235				3 Unchanged High
Q9BZL1	B17gh7	ubiquitin-conjugating enzyme E2 ubiquitin-like 5	4339.474	1980				-1.46E-02 1.07E+00	
P08670	C24cd3	vimentin	5940.876	9100 8720	5279.835 17783.4		1.58E+00		3 Unchanged High 3 Unchanged High
P45880	C18cd3	vollage-dependent anion channe		7280			-5.58E-01		
Q9Y277	C04cd4	voltage-dependent anion channe		2210		2216.75			3 Unchanged High
P17861	O15cd4	X-box binding protein 1	3398.455		2040.224		-7.36E-01		
043570	M04cd3	zinc finger protein 207	3338.117	3170				-7.67E-02	
Q07352	N21ef6	zinc finger protein 36; C3H type-l		5750	4958.224			-8.49E-02	
Q15942	M10cd3	zyxin	2570.21	2070		2387.987			3 Unchanged High
Q16659	B17ef5	mitogen-activated protein kinase		141	100,0111			-3.51E-02	
O75324 Q9BYP8	B05cd3 E16gh8	stannin keratio associated eretain 17.1	131.7242	172			-3.97E-01		
P55291	J01ab4	keratin associated protein 17.1 cadherin 15; M-cadherin (myotub	176.4958	174	100.1011			-2.33E-02	
			.00., 213	128	100.3297	120.0003	-6.62E-01	-0.10E-U1	, orwininged row

P49768	L14ab7	presenilin 1 (Alzhelmer disease :	127.9653	124	100.3504	117.5823	-3.51E-01	-4.04F-02	1 Unchanged Low
P55107	G10ab5	growth differentiation factor 10	114,7206	168	100.6326		-1.89E-01		1 Unchanged Low
P17082	P18cd7	related RAS viral (r-ras) oncoger	433,219	176	100.8465		-2.10E+00		1 Unchanged Low
P23634	N08ab2	ATPase; Ca++ transporting; plas		276	100.9041		-8.30E-01		1 Unchanged Low
P78347	J20ab5	general transcription factor II; i	151.6834	103	100.9514		-5.87E-01		1 Unchanged Low
P09912	F08ef7	Interferon; alpha-inducible protek		1050	101.0749		-8.61E-01		1 Unchanged Low
Q9H2F5	F13gh7	enhancer of polycomb 1	127.3938	115	101.3763		-3.30E-01		1 Unchanged Low
014811	M06cd7	programmed cell death 10	152.3457	132	101.596		-5.85E-01		
060942	A17cd4	RNA guanylyltransferase and 5'-		97.1					1 Unchanged Law
Q9Y5Y3	M18cd7	G prolein-coupled receptor 45	174.3335	161	101.6765 101.8478		-7.98E-01		1 Unchanged Low
P11908	H15ab8			174			-7.75E-01		1 Unchanged Low
Q92851	P18ab3	phosphoribosyl pyrophosphale s	135,919		101.9334		-7.05E-01		1 Unchanged Low
Q9BQE5		caspase 10; apoptosis-related cy		107	101,9435	115.0308			1 Unchanged Low
P55327	M03gh7 D05cd2	apolipoprotein L; 2	109.8196 136.2596	148	102.0533		-1.06E-01	4.26E-01	1 Unchanged Low
P28332		tumor protein D52		112			-4.16E-01		1 Unchanged Low
095861	K03ab2	alcohol dehydrogenase 6 (class '		97.7			-4.79E-01		1 Unchanged Low
	E02cd6	3'(2'); 5'-bisphosphale nucleotida		116			-3.27E-01		1 Unchanged Low
Q9NXJ5 Q15642	A10gh2	hypothetical protein FLJ20208	122.4219	151	102.3038		-2.59E-01	3.05E-01	1 Unchanged Low
	K09ght	thyroid hormone receptor interac		142			-2.41E-01		1 Unchanged Low
094997	009cd7	decidual protein induced by prog		140	102.3137		-7.69E-01		1 Unchanged Low
095453	G02ab8	poly(A)-specific ribonuclease (de		105	102.3217		-4.52E-01		1 Unchanged Low
Q9H9C5	F01gh7	KIAA1453 protein	135.8299	126	102.4347		-4.07E-01		1 Unchanged Low
P10451	B13cd1	secreted phosphoprolein 1 (oster		76.7			-8.94E-01		1 Unchanged Low
P53992	B03cd5	SEC24 related gene family; mem		105	102.5437		-3.89E-01		1 Unchanged Low
Q9NQW1		secretory pathway component Se		158	102.6234		-1.18E-01		1 Unchanged Low
Q01449	A17gh5	myosin light chain 2a	179.1031	170	103.0725		-7.97E-01		1 Unchanged Low
Q9NX46	K12gh2	hypothetical prolein FLJ20446	109.5374	159	103.1083		-8.73E-02		1 Unchanged Low
P53609	A23cd1	protein geranylgeranyltransferas		128	103.1248		-4.41E-01		1 Unchanged Low
Q15029	J16cd4	U5 snRNP-specific protein; 116)		126	103.1503			7.28E-02	1 Unchanged Low
P50458	O23cd5	LIM homeobox protein 2	145,905	120	103.1971		-5.00E-01		1 Unchanged Low
P33764	B24cd1	S100 calcium binding protein A3		204			-5.43E-01		1 Unchanged Low
Q9P289	G03ef3	Mst3 and SOK1-related kinase	133.5225	116	103.4345		-3.68E-01		1 Unchanged Low
P52429	P17cd3	diacylglycerol kinase; epsilon (64		123	103.5015		-3.15E-01		1 Unchanged Low
P54578	H17cd5	ubiquitin specific protease 14 (tR		108	103.6864		-6.33E-01		1 Unchanged Low
Q13219	E24ab8	pregnancy-associated plasma pr		154	103,7948		-1.21E-01		1 Unchanged Low
060547	M12ab5	GDP-mannose 4;6-dehydratase	147.2805	108	103.8004		-5.05E-01		1 Unchanged Low
P23297	117ef6	S100 calcium binding protein A1	145.2558	126	103.9033		-4.83E-01		1 Unchanged Low
Q9H5J8	M01gh6	hypothetical protein MGC5306	216,091	91.7	104.0432		-1.05E+00		1 Unchanged Low
Q14493	108cd4	stem-loop (histone) binding prote		155	104.1318		-2.84E-01		1 Unchanged Low
Q9NZU0	O16cd8	fibronectin leucine rich transmerr		268	104,3033		-1.45E+00		1 Unchanged Low
P80370	B01ef7	delta-like 1 homolog (Drosophila)		80	104.328		-6.18E-01		1 Unchanged Low
Q9NYV4	C23ef3	CDC2-related protein kinase 7	128.5947	107	104,3521		-3.01E-01		1 Unchanged Low
P09016	C23ef5	homeo box D4	173.2254	192	104.3556			1.50E-01	1 Unchanged Low
Q13873	A16ab3	bone morphogenetic protein rece		86.8	104.4206		-5.77E-01		1 Unchanged Low
014878	B22ef4	inosine triphosphatase (nucleosii		120	104.4238		-2.58E-01		1 Unchanged Low
P52848	J01ab6	N-deacetylase/N-sulfotransferasi		84.4	104.4555		-6.70E-01		1 Unchanged Low
Q14848	A05cd5	TNF receptor-associated factor 4		136	104.4604		-2.50E-01		1 Unchanged Low
000204	O04cd2	sulfotransferase family; cytosolic		117	104.4824		-8.31E-01		1 Unchanged Low
P35610	E13gh1	sterol O-acyltransferase (acyl-Cc		88	104,5518		-4.60E-01		1 Unchanged Low
Q04760	LO3ab5	glyoxalase I	132.6871	107	104.5782		-3.43E-01		1 Unchanged Low
P51690	H06ab2	arylsulfatase E (chondrodysplasi	177,023	76.2	104,5865		-7.59E-01		1 Unchanged Low
P26012	C11ef7	integrin; beta 8	110.7396	170	104.6296		-8.19E-02		1 Unchanged Low
Q14978 O00764	K11cd5 D16cd3		117.0048	131	104.685		-1.61E-01	1.61E-01	t Unchanged Low
Q9Y6Q5	P13cd5	pyridoxal (pyridoxine; vitamin B6		140	104.6915		-6.14E-01		1 Unchanged Low
Q910Q5 Q95350	G04cd5		149.3929	107	104.7705		-5.12E-01		1 Unchanged Low
P49748	C13ab2	Homer; neuronal immediate early		126	104.7829		-3.90E-01		1 Unchanged Low
P08100	O24cd1	acyl-Coenzyme A dehydrogenas	121.151	141	104.7897		-2.09E-01		1 Unchanged Low
Q15477	F24cd2	rhodopsin (opsin 2; rod pigment)		116	104.8101		-2.58E-01		1 Unchanged Low
043520	117cd1	superkiller viralicidic activity 2-lik ATPase; Class I; type 88; membi		186	104.8138	131.0636		8.58E-01	1 Unchanged Low
Q13039	P02ef4	ATP-binding cassette; sub-family		91.5	104.8252		-9.26E-01		1 Unchanged Low
015291	L12cd1	solute carrier family 7 (cationic at		80.5 176	104.8585		-7.45E-01 -1.40E+00		1 Unchanged Low 1 Unchanged Low
Q9NR71	L23gh4	milochondrial ceramidase	108.9644	152	104.9361		-5.43E-02		
060884	H10cd4	DnaJ (Hsp40) homolog; subfamil		239	104.9441		-1.08E+00		1 Unchanged Low 1 Unchanged Low
Q9BYE0	J17gh8	halry and enhancer of split 7 (Dr.		108			-2.87E-01		
Q9Y272	L01ef2	RAS; dexamethasone-induced 1					5.29E-02		1 Unchanged Low 1 Unchanged Low
Q9BUR5	M13gh6	hypothetical protein MGC4825	78.2636	239	105.0137		4.24E-01		1 Unchanged Low
060859	A13cd7	neuropathy target esterase	123.9707	117	105.029		-2.39E-01		1 Unchanged Low
Q9GZU1	A20gh4	mucolipin 1	85.32301	243	105.1039	144,4516		1.51E+00	1 Unchanged Low
P19878	B15ab6	neutrophil cytosolic factor 2 (65kl		515	105.1041		-1.56E+00		1 Unchanged Low
Q13277	K12cd2	syntaxin 3A	118.5375	185	105.2511		-1.72E-01		1 Unchanged Low
	B04gh8	reserved	130.849	124	105.2514		-3.14E-01		1 Unchanged Low
P32456	N18ab4	guanylate binding protein 2; Inter		142	105.2611		-5.54E-02		1 Unchanged Low
Q9H4M9	E01cd7	EH-domain containing 1	149.0051	151	105.3893		-5.00E-01		1 Unchanged Low
Q9BTX7	O19gh6	chromosome 20 open reading fra		157	105.3033		-1.37E-01		1 Unchanged Low
Q9NVN2	M02gh3	solute carrier family 4 (anion excl		114	105.466		-1.3/E-01		1 Unchanged Low
Q99758	A17ab2	ATP-binding cassette; sub-family			105.5504		-1.18E+00		1 Unchanged Low
P43694	B06ef6	GATA binding protein 4	107.7539	161	105.5891		-2.93E-02		1 Unchanged Low
Q13825	C07ab3	AU RNA binding protein/encyt-Ct		140	105.5272		-2.93E-02 -2.25E-01		1 Unchanged Low
Q92550	L23ef3	ring finger protein 10	197.4603	112			-9.02E-01		1 Unchanged Low
Q9UJS0	C12ef3	solute carrier family 25; member	119.1806	139			-1.72E-01		1 Unchanged Low
Q99611	O05ef3	selenophosphate synthetase 2	168.8742	128	105,8216		-6.73E-01		1 Unchanged Low
P17275	F08ab6	jun B proto-oncogene	135.9478	94.9	105.0333		-3.60E-01		1 Unchanged Low
Q13477	B09cd4	mucosal vascular addressin cell :		110			-2.71E-01		1 Unchanged Low
Q14692	D09gh1	KIAA0187 gene product	114.5267	131			-1.12E-01		1 Unchanged Low
P49908	D13cd2	selenoprotein P; plasma; 1	140.6083	104			-4.08E-01		1 Unchanged Low
P54803	M24ab3	galactosylceramidase (Krabbe di					-2.99E-01		1 Unchanged Low
		,		•					

P50991	H09cd6	chaperonin containing TCP1; sut	161.4396	148	106,0337	138,5613	-6.06E-01	-1.23E-01	1 Un	changed Low	N
P35548	K11ab7	msh homeo box homolog 2 (Dros	182,4309	124	106,057	137.5011	-7.83E-01	-5.57E-01		changed Low	
P11309	M14ab8	plm-1 ancogene	136.3951	99	106.0757		-3.63E-01			changed Low	
075570	A03cd5	mitochondrial translational releas		118	106.1208		-2.816-01			ichanged Low	
Q9NY61 Q98TY8	001ef3		120,3605	124	106.1404		-1.81E-01	4.36E-02		ichanged Low	
Q9NWS2	D01gh7 F09gh2	hypothetical protein MGC4342 hypothetical protein FLJ20640	139,5304 183,0823	92.9 140	106.2168 106.2384		-3.94E-01	-5.8/E-01 -3.89E-01		changed Low	
000180	H06ab6	potassium channel; subfamily K;	129,4915	122	106,2703			-8.14E-02		ichanged Low ichanged Low	
015382	105ab3	branched chain aminotransferast		95.5	106.3058			-5.19E-01		changed Low	
Q9Y2Z1	D05ef2	CGI-01 protein	156.4563	196	106,3369		-5.57E-01	3.28E-01		changed Low	
Q9Y315	E18ef2	CGI-26 protein	135.7793	111	106.6662		-3.48E-01			changed Low	
Q9Y592	P20ef1	NY-REN-58 antigen	118.6541	121	106.7049	115.3792	-1.53E-01	2.56E-02	1 Un	changed Low	N
Q9NYT0	D20ef3	pleckstrin 2 (mouse) homolog	114.999	153	106.7508		-1.07E-01	4.10E-01		ichanged Low	
Q9Y343	M10ef4	SBBI31 protein	123.0625	120	106,8352			-3.60E-02		changed Low	
Q9BVI4 P30622	G09gh6 B20cd1	hypothetical protein MGC3162	140.5988	123	106,8523			-1.88E-01		changed Low	
Q08345	F08ef5	restin (Reed-Steinberg cell-expre discoidin domain receptor family;		136 264	107.0013			2.15E-01		ichanged Low	
P33176	C16ab7	kinesin family member 58	123.5679	113	107.136			-1.35E-02 -1.33E-01		ichanged Low ichanged Low	
P55273	D14ab4	cyclin-dependent kinase inhibitor		174	107,4234	126,8971	1.08E-01	8.01E-01		changed Low	
Q9NW92	E10gh3	hypothetical protein FLJ10209	177.1237	166	107.454		-7.21E-01			changed Low	
Q9NZC3	K03ef3	membrane interacting protein of I		152	107.465	121.8015	2.21E-02	5.23E-01		changed Low	
P15172	M19ab7	myogenic factor 3	132.7839	110	107.4694	116.7347	-3.05E-01	-2.72E-01	1 Un	changed Low	N
P27144	A06ab2	adenylale kinase 3	118.5733	124	107.5998			6.42E-02		ichanged Low	
O60508 Q14012	A06ef2 B08cd3	pre-mRNA splicing factor 17	150.1968	130	107,6187			-2.09E-01		ichanged Low	
P98179	G12cd1	calcium/calmodulin-dependent ps RNA binding molif protein 3	140.1356	135 93.2	107.6204	118.8965	-8.21E-02			changed Low	
Q13223	E10ab5	BRF1 homolog; subunit of RNA p		115	107.7567	113.6891		-5.89E-01 -3.79E-01		ichanged Low ichanged Low	
P55198	N09ab7	myeloid/lymphoid or mixed-linear		122	107.7762			-5.69E-01		ichanged Low	
P46459	G02ef1	N-ethylmalelmide-sensitive factor		95.7	107.8044	114,1635		-5.37E-01		ichanged Low	
P34969	121 ef1	5-hydroxytryptamine (serotonin) ı	218.7666	181	107.8181	169.3507	-1.02E+00	-2.70E-01		ichanged Low	
Q14667	K02gh1	KIAA0100 gene product	105.9471	222	107.8612	145.1883	2.58E-02	1.07E+00		changed Low	
Q92830	J14cd8	GCN5 general control of amino-a		106				-2.83E-01		schanged Low	
O43252	P14cd2	RNA binding motif protein; Y chro		153	107,9125		-5.87E-02			changed Low	
Q9NS93	J01cd5 G09ef3	3'-phosphoadenosine 5'-phospho seven transmembrane protein Til		188 237	107.9154			-5.02E-02		ichanged Lov	
060616	E20ab6	core-binding factor; runt domain;	489.0924	48.1	107.9483 107.9969		-3.55E-01	7.76E-01 -3.35E+00		ichanged Low ichanged Low	
P02261	D19cd4	H2A histone family; member C	145.0821	84.1	108.023			-7.87E-01		ichanged Low	
Q14701	F18ef7	DNA cross-link repair 1A (PSO2	119.2486	307	108.208		-1.40E-01	1.37E+00		changed Low	
Q9P025	A04ef8	HSPC135 protein	160.3109	135	108,2493	134.6437	-5.67E-01	-2.44E-01		changed Low	
P06702	O12ef7	S100 calcium binding protein A9	144.7319	163	108.2576	138,5985		1.70E-01	1 Un	changed Low	N
Q9Y5Q9	P05cd7	general transcription factor IIIC; r	138,644	220	108.3524		-3.56E-01	6.68E-01		changed Low	
P19075 O60905	O20cd2 B16cd4	transmembrane 4 superfamily me sperm associated antigen 9	129.5468	103	108.6355		-2.54E-01			changed Low	
Q13322	F24ef5	growth factor receptor-bound pro		139 115	108.6913 108.7182		-2.73E-02	3.27E-01 -7.64E-02		ichanged Low ichanged Low	
Q9UQ09	L01gh1	KIAA0470 gene product	190.3667	103	108,9005			-8.89E-01		changed Low	
P57053	J20gh6	H2B histone family; member S	113.6057	199	108.9238	140,5964	-6.07E-02			changed Low	
Q92876	J03ab8	kallikrein 6 (neurosin; zyme)	197.2979	130	108.9308	145.5667	-8.57E-01	-5.97E-01		changed Low	
P09960	F21ab6	leukotriene A4 hydrolase	96.36818	185	109,0526	130.2822	1.78E-01		1 Un	changed Low	N
Q9HBH1	002gh5	peptide deformylase-like protein	138.0653	91				-6.01E-01		changed Low	
P55082 Q9NVH6	L15ab7 B01gh3	microfibrillar-associated protein :		124	109.134		-1.54E-01			ichanged Low	
O95249	K10cd5	trimethyllysine hydroxylase; epsil golgi SNAP receptor complex me	107.5878	81.6 143	109.1442 109.1657	121,828 119,983	2.10E-02	-1,10E+00		changed Low	
Q13891	M15ab6	basic transcription factor 3; like 2		120	109.1942			4.12E-01 -8.18E-02		ichanged Low ichanged Low	
Q9BZM5	H05gh7	UL16 binding protein 2	133.6731	245	109.2193	162.5485				changed Low	
O94913	O20ef2	PCF11p homolog	205.6752	80.3	109.2409			-1.36E+00		changed Low	
Q12987	E03ef4	acidic 82 kDa protein mRNA	132.1136	117	109.4318	119.3623	-2.72E-01	-1.81E-01	1 Un	changed Low	N
Q92499	D05ab5	DEAD/H (Asp-Glu-Ala-Asp/His) t		124	109.4356			-5.13E-01		ichanged Low	
Q13686 P43250	B19ab2 108ab4	alkylation repair; alkB homolog	105.5169	143	109.4797	119.4385				changed Low	
Q9UMX0	L03ef4	G protein-coupled receptor kinas ubiquilin 1	122.0111	113 141	109.4844 109.5522		-1.37E-01 -1.55E-01			ichanged Low	
O94905	G12cd7	chromosome 8 open reading fran		107	109.6202			-2.57E-01		ichanged Low ichanged Low	
Q9UHK0	K01cd8		157.4757	87.6	109.6296	118.2396		-8.46E-01		changed Low	
P25189	D01ab6	myelin protein zero (Charcot-Mar	110.5502	132	109.6859	117,3068	-1.13E-02	2.52E-01		changed Low	
Q10567	M11ab2	adaptor-related protein complex	208.9283	187	109.8204	168,4738	-9.28E-01	-1.62E-01	1 Un	nchanged Low	N
O15230	D18cd8	laminin; alpha 5	111.864	141				3.35E-01	1 Un	ichanged Low	N
O60668	J10ab6	Karyopherin alpha 1 (importin alp		169	109.9831			-7.70E-02		ichanged Lov	
P11686	M03cd2 J02cd1	TAF2 RNA polymerase II; TATA surfactant; pulmonary-associatec		111				-1.56E-01 -2.19E-01		ichanged Low	
Q9NWB6		hypothetical protein FLJ10154	174.7789	117 75	110.0847			-2.19E-01 -1.22E+00		ichanged Low ichanged Low	
Q9H028	E22gh8	histone deacetylase 10	107,5034	138				3.57E-01		ichanged Low	
Q9UMZ2	O22cd7	AP1 gamma subunit binding prot		119				-1,32E-02		changed Low	
P24752	C15ab2	acelyl-Coenzyme A acetyltransfe	216.1409	114	110.2562			-9.23E-01		changed Low	
P11230	B06ab3	cholinergic receptor; nicotinic; be		151	110.2999			5.11E-01	1 Un	ichanged Low	N
014977	O18ef2	omithine decarboxylase antizym		129				1.06E-01		changed Low	
P56937	E15ef5 N06ef2	T cell receptor beta locus hydroxysteroid (17-beta) dehydro	139.7091	844				2.59E+00		nchanged Low	
O75299	B21ab7	kinesin family member C3	113.15	74.5 128				-1.61E+00 1.78E-01		changed Low	
P16219	C09ab2	acyl-Coenzyme A dehydrogenas			110.3535	117.1803		3.16E-01		ichanged Low ichanged Low	
Q9NZ48	J20gh3	uncharacterized hemalopoietic st		189	110.3995			7.46E-01		ichanged Low	
Q9P163	L10gh4	hypothetical protein PRO2521	101.1317	159	110.4184	123,4514	1.27E-01	6.51E-01		ichanged Low	
Q9BVT8	107gh8	hypothetical protein MGC5442	135.5302	137	110.5044	127.7419	-2.95E-01	1.76E-02	1 Um	nchanged Low	
Q99933	E02ef6	BCL2-associated alhanogene	119.7332	115				-5.98E-02		nchanged Low	
O14645 Q9BRR9	M12cd3 E01gh8	dynein; axonemal; light intermedi Rho GTPase activating protein 9		207				3.31E-01		nchanged Low	
Q9Y3E7	J19ef2	CGI-149 protein	158.4397	115 202				-7.10E-02 3.54E-01		nchanged Low	
P46976	C09ab6	glycogenin	127.6502	117				-1.21E-01		ichanged Lov ichanged Lov	
									. 01		

075928	C01cd5	Protein inhibitor of activated STA	98.84393	173	110.7286	127.4786	1.64E-01	8.06E-01		
Q9NX63	118gh2	hypothetical protein FLJ20420	123.3725	159	110.8068		-1.55E-01			Unchanged Low
Q9UBS0	120cd2	ribosomal protein S6 kinase; 70k	186.953	155						Unchanged Low
Q15185	F18cd6	unactive progesterone receptor;			110.8188			-2.71E-01		Unchanged Low
				72,3	110.8534			-1.23E+00	1	Unchanged Low
060516	J18cd3	eukaryotic translation initiation fa		147	110.9085			4.97E-01		Unchanged Low
Q16651	J01ab8	protease; serine; 8 (prostasin)	334,0498	125	110.9253			-1.41E+00		Unchanged Low
P48788	E22cd2	troponin I; skeletal; fast	121,3602	110	110,9357				1	Unchanged Low
Q01968	N06ab7	oculocerebrorenal syndrome of L		95.9	110.9507	125.3025	-6.07E-01	-8.17E-01	1	Unchanged Low
Q9H6Z6	L11gh7	hypothetical protein FLJ21628	115.1546	126	111.0337	117.2667	-5.26E-02	1.25E-01	1	
Q9NQ55	B01gh4	peter pan homolog (Drosophila)	122.449	120	111,0627	117.676	-1.41E-01	-3.50E-02	1	
095359	A24gh1	transforming; acidic coiled-coil ct	108.2194	139	111,0999	119.4624	3.79E-02	3.82E-01		Unchanged Low
P01189	115ab8	prooplomelanocortin (adrenocort	145.0634	90.4	111,1389				1	
Q9Y525	G13cd8	v-mat musculoaponeurolic fibros	119,023	141	111,1934				4	
Q9UHC9	L09cd8	NPC1 (Niemann-Pick disease; ty		138	111.3741		4.79E-02	3.53E-01	i	
Q9HBT2	P14gh7	hypothetical protein PP1057	147,5097	98	111,4385					Unchanged Low
000219	M22ef1	hyaluronan synthase 3	224,5087	89	111.4539			-1.34E+00		
P22735	M16cd1	transglutaminase 1 (K polypeptid		91	111.5589				1	
015388	A11ab4	sialic acid binding lg-like lectin 6	105,2847	147					1	
095389	J01cd4	WNT1 inducible signaling pathwr			111.6016 111.6123			4.82E-01		Unchanged Low
Q06190	D21ab8	protein phosphatase 2 (formerly:		94						Unchanged Low
P50226	C03cd3			102	111.8192					Unchanged Low
		sulfotransferase family; cytosolic		101	111.8328					Unchanged Low
Q16632	N10ab5	general transcription factor IIIC; p		116	111,85				1	Unchanged Low
P54762	A24el7	EphB1	107.3397		111.8669				1	Unchanged Low
Q15269	E15cd1	PWP2 periodic tryptophan protei		107	111.8713		-1.61E-01	-2.25E-01	1	Unchanged Low
014495	H10cd3	phosphalidic acid phosphalase to	120,761	109	111.9087	113.9808 -	-1.10E-01	-1.44E-01	1	
Q9P0U0	N24ef3	PC326 protein	152,5639	101	111.9543	121.9267	-4.47E-01	-5.91E-01	1	Unchanged Low
P50120	J02ab8	relinol binding protein 2; cellular	231.806	217	111.9639	187.0028 -	1.05E+00	-9.36E-02	1	Unchanged Low
Q9UHR0	D08gh1	G protein-coupled receptor kinas	111,83	151	111.9778	124.9147	1.91È-03	4.33E-01	1	
Q13977	A12ef1	cerebellar degeneration-related ;		180	112,0477	132.4319			1	
P51692	C18cd8	signal transducer and activator o	126,7779	100	112.0645				i	
P51164	N12ab2	ATPase; H+/K+ exchanging; beta	136.4695	112	112.2007				1	
Q9NXH3	C02gh2	protein phosphatase 1; regulator	113,3081	127	112.2521	117.6836		1.70E-01	i	
	C04gh7	ER to nucleus signalling 2	112,9203	125	112,2701	116.8658		1.51E-01		Unchanged Low
P16435	G16ef6	P450 (cytochrome) oxidoreducta:		132	112,2899		2.71E-02	2.62E-01		
	E05gh6	hypothetical protein MGC5338	116,1511	137	112.3522	121.7814				Unchanged Low
Q02080	L03ab7	MADS box transcription enhance		100				2.36E-01		Unchanged Low
Q15334	J06ab7	lethal glant larvae homolog 1 (Dr			112.3745	117.0502 -				Unchanged Low
P10636	J13ab7	microtubule-associated protein to			112.3893	115.3049 •				Unchanged Low
Q9Y4J6				_	112.4092			2.94E-01		Unchanged Low
	A01gh2	zinc linger protein 6 (CMPX1)	138.8767		112.4376	117.8247 -			1	Unchanged Low
Q9UHY7	P18gh4	E-1 enzyme	108,9217	134	112.5485		4.73E-02	3.00E-01	1	Unchanged Low
Q14147	K04gh1	DEAD/H (Asp-Glu-Ala-Asp/His) t		119	112,7866	116.4348 -	-5.95E-02	1.76E-02	1	Unchanged Low
O95880	N06gh4	KIAA1513 protein	168,2097	104	112,9115	128,5046 -	5.75E-01	-6.88E-01	1	Unchanged Low
Q15013	J23ef3	gene predicted from cDNA with a	104,556	198	113.0807	138.5918	1.13E-01	9.22E-01	1	Unchanged Low
Q9P1E2	E01gh4	hypothetical protein PRO2219	138.6039	107	113.1066	119.6869 -	2.93E-01	-3.69E-01		Unchanged Low
P07203	H23ef1	glutathione peroxidase 1	120.5127	114	113.1788	115.8519 -	9.06E-02	-8.19E-02		Unchanged Low
Q13206	M07ab5	DEAD/H (Asp-Glu-Ala-Asp/His) t		92.5	113.1804	115.4153 -	3.12E-01	-6.03E-01	1	
O15085	L15ef3	Rho guanine nucleotide exchang		111	113.2492	115.3055 -	1.05E-01	-1.37E-01	1	
P48426	C07cd1	phosphatidylinositol-4-phosphate	232.9191	163	113.2533	169.6778 -1			1	
Q9NRQ2	K20gh4	phospholipid scramblase 4	126,2397	116	113.2849	118.4782 -			i	goo bon
043171	B24cd3	CDC14 cell division cycle 14 hon	126.784	113	113.382	117.8862 -			i	
Q9HB90	M02gh5	Rag C protein	132.9037		113.4299	127.5658 -		3.71E-02	i	
Q9H733	121gh5	single lg IL-1 R-related molecule	113,5449	137	113,4753	121.334 -		2.71E-01	i	
P43353	C06ab2		141,7651		113.5244	118.734 -		-4.90E-01		
Q15053	G14gh1	KIAA0040 gene product	111.1705		113.5775					Unchanged Low
P22301	H01e/7	interleukin 10	174.8595				3.09E-02	3.91E-01		
Q9BW47	N12gh5	chromosome 20 open reading fra			113.5877	132.021 -			1	
P21917	M03ab6	dopamine receptor D4			113.6231	131.1841 -		2.92E-01	1	Unchanged Low
Q9Y6K5	M19cd1	2-5'-oligoadenylate synthetase 3	158,6974		113.7899	141.2355		-6.96E-02	1	Unchanged Low
					113.9993	127.6931 -		1.56E-01	1	
Q9Y2A4	N22cd5	Kruppel-type zinc finger (C2H2)	136.0849		114.1396	127.5854 -		-3.82E-02	1	
Q16880	A16cd3		112.9309		114.1612	119.9407	1.56E-02	2.33E-01	1	Unchanged Low
O95747	D09cd5	oxidative-stress responsive 1	113.39		114.2384		1.08E-02	1.12E-01	1	Unchanged Low
P25106	L12ef4	G protein-coupled receptor	100.7123				1.82E-01	5.42E-01	1	Unchanged Low
	H23ab6		102.6971	145	114,3219	120.6924	1.55E-01	4.98E-01	1	
P45984	L18ef5		139.4168	88	114.3405	113.914 -	2.86E-01	-6.64E-01	1	
Q9NPF7	103ef3	interleukin 23; alpha subunit p19		182	114.4153	131.5891	2.11E-01	8.77E-01	1	Unchanged Low
Q9UH94	L07cd8	prolactin regulatory element bind				120.3231 -				Unchanged Low
Q16559	F05cd2	T-cell acute lymphocytic leukemi:	131.4974	111	114.5482	118.9991 -	1.99E-01	-2.45E-01		Unchanged Low
Q9UGH0	C02cd8	solute carrier family 17 (anion/su	98.73308		114.6616		2.16E-01	6.75E-01		Unchanged Low
Q15544	J13cd2	TAF11 RNA polymerase II; TATA	117.4634		114.6792	118.4317		6.82E-02		Unchanged Low
000628	N24ab7	peroxisomal biogenesis factor 7	156,16		114.7281	122.588 ~				Unchanged Low
043241	F07ab5	golgi autoantigen; golgin subfam			114.8381	114.422				
Q9Y6N1	M14ab4	COX11 homolog; cylochrome c c				114.0678 -				Unchanged Low
Q9UHX5	001ef4	HT002 protein; hypertension-rela								Unchanged Low
Q01664	A02cd2	transcription factor AP-4 (activati				122.8348		6.60E-01		Unchanged Low
Q9BUI1	003gh6	leukocyte receptor cluster (LRC)			115.0377	150.0321 -1				Unchanged Low
Q9HBL5	M17gh4	AD023 protein		125	115.039		3.55E-02	1.53E-01		Unchanged Low
			103.9999				1.46E-01	3.93E-01		Unchanged Low
Q9NXV2	N08gh2	hypothetical protein FLJ20040	126.6647			129.6462 -		2.16E-01		Unchanged Low
Q15424	D08cd1		134.9613			114.2781 -2				Unchanged Low
P26440	F06ab6	isovaleryi Coenzyme A dehydrog				136.2534 -				Unchanged Low
O60264	J11cd3	SWI/SNF related; matrix associa:		132	115.2498	117.7402	1.24E-01	3.22E-01		Unchanged Low
Q15286	K07cd7	RAB35; member RAS oncogene		183	115.3443		2.62E-01	4.06E-01		Unchanged Low
Q9UGK0	110ef4		303.2745			196.7868 -1		-8.21E-01		Unchanged Low
Q92980	J06cd5	pulative protein similar to nessy (142.089		115.4166	146.2003 -		3.50E-01		Unchanged Low
Q99726	M06cd3	solute carrier family 30 (zinc tran	112.8008				3.39E-02			Unchanged Low
P01111	O17ab8		123.6283			115.8529 -				Unchanged Low
Q92574	G13cd3		109.8519			117.8303				Unchanged Low
									•	e-mingen cow

P16106	C01gh7		134.9364	118	115 869	122 8016	-2.20E-01	4 005 04		
Q9NWX		hypothetical protein FLJ20548	118.1581	112			-2.67E-02			1 Unchanged Low
Q9Y6B7			· 217.8572	170		168.0108	9.09E-01	-7.20E-02		1 Unchanged Low
Q9Y4F1	J04cd5	FERM; RhoGEF (ARHGEF) and	108.6215	130		118.0819	9.52E-02	2.55E-01		1 Unchanged Low
Q06710	M14cd3	paired box gene 8	120.1357	129			-4.93E-02			1 Unchanged Low
Q9H6N1	J17gh5	hypothetical protein FLJ22056	214.4909	242			-8.84E-01			1 Unchanged Low
Q9NSY1			# 111.9034	123						1 Unchanged Low
Q14106	H20cd6	transducer of ERBB2; 2	117.9634	134						1 Unchanged Low
Q9UK11	B02ef4	zinc finger protein 223	119.1221	133			-3.47E-02			Unchanged Low
P08069	017ef6	insulin-like growth factor 1 rece	pl 106.3434	178	118,3103	133,4883		7.42E-01		Unchanged Low
Q9H6F5	121gh6	hypothetical protein MGC2574	120.5097	112			-5.11E-02	-1 10E-01		Unchanged Low
Q14373	O04ab3		na 157.7476	231			-4.39E-01			Unchanged Low
P20393	G04ef5	nuclear receptor subfamily 1; gr	a 107.1138	138	116,4228					I Unchanged Low I Unchanged Low
P36915	O01ab5	guanine nucleolide binding prot	e 120.9306	111	116,4819		-5.41E-02			Unchanged Low
095109 Q9NVS2	G02cd8	chromosome 20 open reading fr	z 143.5728	103	116.5321	121.0452	-3.01E-01	-4.79E-01		Unchanged Low
		mitochondrial ribosomal protein	£ 117.9573	124	116.6011	119,5497	-1.67E-02	7.31F-02		Unchanged Low
Q92950	B08cd6	CUG triplet repeat; RNA binding	109.8876	129	116.7366	118.6125	8.72E-02	2.34E-01		Unchanged Low
Q14977	C04ab7	SP110 nuclear body protein	107.0969	158	116.8973	127.3783		5.62E-01		Unchanged Low
P16106 Q9Y3C1	O04gh6	H3 histone family; member A	94.76745	159	116.9796	123,5385		7.45E-01		Unchanged Low
P37287	N18ef2	hypothetical protein HSPC111	145.1935	164	117.0185	141.934	-3.11E-01	1.72E-01		Unchanged Low
Q9UIG5	C11ef1	phosphatidylinositol glycan; clas		109	117.2467		-1.77E-01			Unchanged Low
Q9UNE2	G05ef4	SEEK1 protein	102.5149	148		122.443	1.94E-01			Unchanged Low
Q14123	G15cd6 A17cd1	rabphilin 3A-like (without C2 dor	r 110.3023	123				1.62E-01		Unchanged Low
Q16206	P03ab2	phosphodiesterase 1C; calmodu	121.3078	132			-4.75E-02	1.21E-01		Unchanged Low
Q9NWT1		cytosolic ovarlan carcinoma anti		157		130,1671	1.15E-02	4.26E-01		Unchanged Low
Q12756	C05ab3	hypothelical protein FLJ20624	116.0517	116		116.4708		-2.15E-03		Unchanged Low
P28288	B10ab8	axonal transport of synaptic vest	138.4398	232			-2.36E-01	7.47E-01		Unchanged Low
P05198	G08ab5	ATP-blnding cassette; sub-family	140.4464	95.9			-2.56E-01	-5.51E-01		Unchanged Low
Q9H0V9	D08gh7	eukaryotic translation initiation fa hypothetical protein DKFZp564L	115.50/4	119				3.77E-02		Unchanged Low
Q92984	M10ef1	Interferon-induced protein 35	258.0737	101	117.6689		-1.31E-01	-3.47E-01	1	
095498	A13cd5	vanin 2		221	117.8182		-1.13E+00		1	Unchanged Low
Q9H1E1	J01gh8	ribonuclease 7	109.7731 109.332	123	117.8273	118.8572		1.64E-01	1	
Q9NV88	H01gh3	hypothetical protein FLJ10871	108.5849	131	117.8377	119.4295	1.08E-01	2.62E-01	1	
Q9Y310	F06cd6	SnRNP assembly defective 1 ho	119 9305	125 113	117.8411	117.172	1.186-01	2.04E-01	1	
P48382	O06cd1	regulatory factor X; 5 (influences	191 5142	122	110.0003	117.0556	-2.52E-02	-8.10E-02	1	
P52738	K04cd3	zinc finger protein 140 (clone pH	136 6586	154	118,1501		-6.98E-01		1	
Q9NRW1		RAB6B; member RAS oncogene	100.578	179	118.1659		-2.10E-01	1.71E-01	1	
O15298	A02ab3	basic leucine zipper nuclear fact	266.9887	110	118,2062		2.32E-01 -1.18E+00	8.29E-01		Unchanged Low
Q9NUN5	D14gh3	hypothetical protein FLJ11240	114.5392	116	118.2691					Unchanged Low
Q07699	L03cd1	sodium channel; voltage-gated; t	116.5431	113		115.8757	4.62E-02 2.19E-02			Unchanged Low
P27701	F10ab6	kangai 1 (suppression of tumorig	116.2761	108		114.2624	2.608-02		1	Unchanged Low
Q9UER5	O02ef3	TNF-induced protein	104.9755	131		118.2694	1.74E-01	3.24E-01	1	Unchanged Low
QBTAC6	124gh7	chromosome 11 open reading fra	117.4257	138	118,5887	124,5089	1.42E-02		1	Unchanged Low
Q9NRW9	K16gh4	angiotensin II; type I receptor-ass	136.6357	105	118.6777		-2.03E-01		1	Unchanged Low
095147	O15cd7	dual specificity phosphatase 14	181.9968	200		166.9415	-6.17E-01	1.37E-01	i	Unchanged Low
Q13895	A19ef7	byslin-like	108.0232	124		116.9004	1.37E-01	1.98E-01	i	Unchanged Low Unchanged Low
P31645 P51511	N03ef5	solute carrier family 6 (neurotran-	114.6464	189		140.7231	5.19E-02		i	Unchanged Low
Q15154	N17ef7	matrix metalloproteinase 15 (mer		125		122,1739	-4.82E-02	2.16E-02	i	Unchanged Low
Q00007	Officet	pericentriolar material 1	127.8378	139		128.7076		1.25E-01	i	Unchanged Low
P55822	E17ef6 L08cd2	protein phosphatase 2 (formerly:	108.364	123	118.9838	116,7703	1.35E-01	1.82E-01	i	Unchanged Low
Q14442	L20ab8	SH3 domain binding glutamic ad	131.7189	133	119.0263	128.0275	-1.46E-01	1.76E-02		Unchanged Low
P54792	L04ab4	phosphatidylinositol glycan; class		114	119.0306	118.0476	-2.44E-02	-8.61E-02		Unchanged Low
P42336	A03ef6	dishevelled; dsh homolog 1 (Dro:	126,677	113	119.067	119.4338	-8.94E-02	-1.70E-01	1	Unchanged Low
	G15ab3	phosphoinosllide-3-kinase; catal: BarH-like homeobox 2			119.1581	121.01	2.37E-01	4.98E-01	1	Unchanged Low
	M04ef3	solute carrier family 7; (cationic a	99.5713			132.6879	2.59E-01		1	Unchanged Low
	H01ef6	hislidine decarboxylase			119.3079	120.9226	-5.44E-01 -	1.32E+00		Unchanged Low
	B16ef3	N-acetylglucosamine-phosphate	111.9277		119.3155	116.6743	9.22E-02	8.57E-02		Unchanged Low
	A14ab2	aminolevulinate; delta-; synthase	202 4470		119.3184	118.8198	-6.50E-02	-1.52E-01	1	Unchanged Low
	G10ef2	CGI-94 protein	108.7497				-7.60E-01			Unchanged Low
Q9H5J4	105gh6	long-chain fatty-acyl elongase	135.0119				1.36E-01	6.16E-01		Unchanged Low
P16083	E15ab8	NAD(P)H dehydrogenase; quinor	154 7986			113.8461	-1.76E-01	-6.34E-01		Unchanged Low
Q99819	D02ab2	Rho GDP dissociation inhibitor (I	125 7962		119.5724 119.5853	124.3290	-3.73E-01	-6.50E-01		Unchanged Low
	F06cd3	protein kinase; interferon-inducib	100.3086		119.6257	110 0057	-7.30E-02 ·			Unchanged Low
	F12ef7	serine/arginine repetitive matrix 2	125.19	96.6	440 740	***		4.47E-01		Unchanged Low
Q9HBH5	K09gh4	PAN2 protein	102.0601		119.7321		-6.45E-02 - 2.30E-01			Unchanged Low
	C22ab8	purinergic receptor P2Y; G-prote	131.9219				-1.39E-01	4.402-01		Unchanged Low
	D01cd6	RAN binding protein 8	114.6925				6.43E-02			Unchanged Low
	C17cd7	mucosa associated lymphoid tiss	114.4769		119.9403		6.73E-02			Unchanged Low
	E22cd5	mitogen-activated protein kinase	92.42403	156			3.76E-01			Unchanged Low
	J01ef5	sarcoma amplified sequence	114.9397				6.17E-02 -			Unchanged Low
	D07cd2	bromodomain containing 2	116.6707				4.08E-02			Unchanged Low
	H21ef7	ephrin-A1	298.0458			392.8203 -	1.31E+00	1.35E+00		Inchanged Low
	P20gh8	hypothetical protein MGC4171	129.5733		120,4428	118.9065	1.05E-01 -	2 80E-01		Unchanged Low
	L20ef4		105.4425		120.5386	118.2227		2.87E-01		Unchanged Low
	G15gh8	hypothetical protein MGC10812	148.2279	273		180.6391 -		8.82E-01		Unchanged Low
	F04ef5	transforming growth factor; beta	115.7567	108			6.00E-02 -	9.83F-02		Unchanged Low
	E08ef3	YME1-like 1 (S. cerevislae)	256.0314		120,7227	191.0958 -	1.08E+00 -	3.82E-01		Jnchanged Low Jnchanged Low
	G11ab3	BRCA1 associated protein-1 (ubl	101.36					6.72E-01	4	Jochanged Low
	305gh2	hypothetical protein FLJ20604	99.76081					6.26E-01	4 1	Inchanged Low
	N14ab8	Rab geranylgeranyltransferase; a			120.7706 2	205.3234 •	8.83E-01	2.91E-01		Inchanged Low
		apolipoprotein L; 3	106.592	130	120.798	119.152B	1.80E-01	2.87E-01		Inchanged Low
		dual specificity phosphatase 8	146.3963		120.7987	131.9045 -	2.77E-01 -	1.88E-01		Inchanged Low
		spectrin; beta; erythrocytic (inclu-	112.72	114	120.8055	16.0084	9.99E-02	2.26E-02		Inchanged Low
7500B1	218ef1	putative glycolipid transfer protei	99.78536	143 1	120.8708		2.77E-01			Inchanged Low

P05423	A20ab3	BN51 (BHK21) temperature sens	131 8122	95.5	120.9201	116.0734	-1.24E-01	-4.65E-01	1 Unchanged Low
Q14338	117ab6	fucosyltransferase 2 (secretor str			120.9483		2.78E-01	8.63E-01	1 Unchanged Low
Q9BYI8	A22gh8		90.11911		121.0209		4.25E-01	8.67E-01	1 Unchanged Low
	D01ab8		216.3154	105	121.029		-8.38E-01		1 Unchanged Low
Q06203 Q9NZ33	P10gh4		113.8613		121.0508	132,4428		5.12E-01	1 Unchanged Low
P04196	O10ab6		116.1305		121,1163	117.5882	6.06E-02	-7.63E-03	1 Unchanged Low
O96029	F03cd7		201.3579		121,2888	138.7174			1 Unchanged Low
Q01196	116ab5		441,1217		121,3316	232.9078 -			1 Unchanged Low
Q9UHL4	J17cd8		180.3445	191	121,351		-5.72E-01		1 Unchanged Low
P55211	P16ab3	caspase 9; apoptosis-related cys					9,62E-02		1 Unchanged Low
P21283	A07ab3		288,6034		121.3828	232.7446 -			1 Unchanged Low
043422	P02ab8	protein-kinase; Interferon-inducit		100	121.394	117.7416			1 Unchanged Low
P78406	L01cd3	RAE1 RNA export 1 homolog (S.			121,4625	121,9098			1 Unchanged Low
P43362	J18ab7		99.05783		121.5326	119.97	2.95E-01	4.92E-01	1 Unchanged Low
Q9H693	E12gh6	hypothetical protein FLJ22477	171,7699		121.6097		-4,98E-01		1 Unchanged Low
Q9UJB3	A11cd8	2-hydroxyphytanoyl-CoA lyase	99.45467		121.6399		2.91E-01	4.99E-01	1 Unchanged Low
Q05048	115ab4	deavage stimulation factor, 3° pn			121.6764		-2.92E-02		1 Unchanged Low
P19784	D11ab4		94,68573		121.7819	122.6579	3.63E-01		1 Unchanged Low
Q9BUX8	D06gh6	homolog of yeast ribosome blogs			121.8214	155,6701		8.56E-02	1 Unchanged Low
Q9Y5P8	D23cd8		166.4572		121,8219	125.8828			1 Unchanged Low
Q9Y3E1	F16ef1		97.00713		121.9163	122.172	3,30E-01	6.05E-01	1 Unchanged Low
P40426	007cd1		122.6888		121,9253		-9.01E-03		1 Unchanged Low
P11217	K01cd1	phosphorylase; glycogen; muscle		110	122,008			-1.52E+00	1 Unchanged Low
Q16563	D18cd2	synaptophysin-like protein	97.62823		122.0944				1 Unchanged Low
O95698	L07ab6	inhibitor of growth family; membe			122.1358	119.6732	2.57E-01		1 Unchanged Low
P19447	K24ef6	excision repair cross-complemen			122.1425	121.7269	3.56E-01		1 Unchanged Low
Q9UK39	D08cd7	CCR4 carbon catabolite repressi			122.1589		-5.09E-01		1 Unchanged Low
Q15785	E23cd7	translocase of outer mitochondria		144	122.1776			-1.04E+00	1 Unchanged Low
Q99571	C14ab8	purinergic receptor P2X; ligand-ç		410	122.3053			-2.96E-02	1 Unchanged Low
Q9BTY2	E24gh8		131.9449		122,3524			-5.28E-01	1 Unchanged Low
Q9P0P1	D12eft	hypothetical protein FLJ10769	147.9172	191	122,4861		-2.72E-01		1 Unchanged Low
Q9Y324	P01ef1	CGI-35 protein	129.1118	97	122,5531			-4.13E-01	1 Unchanged Low
P37286	017cd3	ubiquilin-conjugating enzyme E2		165	122,5669		3,10E-01		1 Unchanged Low
Q00577	007ef5	purine-rich element binding prote		123	122.5705	120.658	7.91E-02		1 Unchanged Low
O75152	L21gh1	KIAA0663 gene product	101,4464	140	122.5789		2.73E-01		1 Unchanged Low
P23610	H06cd7	coagulation factor VIII-associater	164.911	103	122,5902			-6.73E-01	1 Unchanged Low
O75438	M20cd5	vacuolar protein sorting 26 (yeas		108	122,6522	116,2363		-1.30E-01	1 Unchanged Low
Q9HA40	B23gh5	hypothetical protein FLJ12287 si		126	122.666		2.29E-01		1 Unchanged Low
Q9NX62	120gh2	hypothetical protein FLJ20421	103,7764	136	122,701		2,42E-01		1 Unchanged Low
O76062	E04cd2	transmembrane 7 superfamily me		215	122,7645			-4.89E-01	1 Unchanged Low
Q16445	H02ab3	gamma-aminobutyric acid (GABA	150,641	87.5	122,7829			-7.83E-01	1 Unchanged Low
Q08462	G11ef6	adenylate cyclase 2 (brain)	107.541	146	122.8502		1.92E-01		1 Unchanged Low
P78560	A22ef6	CASP2 and RIPK1 domain conta		153	122.8627	128,1152	1.80E-01		1 Unchanged Low
P19634	N17ef5	solute carrier family 9 (sodium/hy		99	122,898			-2.66E-01	1 Unchanged Low
P28370	N12cd1	SWI/SNF related; matrix associa-	115.471	106	122.9184	114,9572	9.02E-02	-1.17E-01	1 Unchanged Low
Q92503	H06cd1	SEC14-like 1 (S. cerevislae)	151,5939	151	122,9424	141.9225	-3,02E-01	-3.46E-03	1 Unchanged Low
Q9UL02	H24ef1	L-kynurenine/alpha-aminoadipati	105.6951	124	123,0487	117,4294	2.19E-01	2.25E-01	1 Unchanged Low
Q9Y5T5	J13cd6	ubiquitin specific prolease 16	117,1056	120	123,0646	120.0007	7.16E-02	3.32E-02	1 Unchanged Low
	G03gh8	protocadherin 15	153.0978	126	123.1548	134,1729	-3.14E-01	-2.78E-01	1 Unchanged Low
P15848	H02ab2	arylsulfatase B	149.4791	95.6	123.1651	122.7499	-2.79E-01	-6.45E-01	1 Unchanged Low
Q9UP66	P24ab6	low density lipoprotein receptor-r	114.2435	124	123,2083	120.4437	1.09E-01	1.17E-01	1 Unchanged Low
O75160	P07gh1	KIAA0672 gene product	121.9398	117	123.2554	120.8501	1.55E-02	-5.53E-02	1 Unchanged Low
P78385	L14ab6	keratin; hair; basic; 3	100.703	135	123.2569	119.6793	2.92E-01	4.24E-01	1 Unchanged Low
O94967	B21ef8	KIAA0893 protein	92.15324	158	123.2707	124.6247	4.20E-01		1 Unchanged Low
O95719	K01cd5	xenotropic and polytropic retrovir	108.0877	122	123.3497	117.7043	1.91E-01	1.71E-01	1 Unchanged Low
P48147	F09ab8	prolyl endopeptidase	130.0641	129	123.3564			2 -7.45E-03	1 Unchanged Low
P20231	B15gh7	tryptase beta 2	106.6865	130	123,4905				1 Unchanged Low
O95299	G02ab7	NADH dehydrogenase (ubiquino	107,385	129	123,5991				1 Unchanged Low
075033	118gh1	KIAA0445 gene product	121.5196	122	123.624			9.32E-03	1 Unchanged Low
P51668	D03cd4	ubiquitin-conjugating enzyme E2		137	123,637				1 Unchanged Low
P28289	E08cd2	tropomodulin	303.877	272	123.6484			-1.62E-01	1 Unchanged Low
Q15631	O12ef6	translin	173,134	152	123.6887			1 -1.88E-01	1 Unchanged Low
P49888	F03cd2	sulfotransferase; estrogen-prefer		95.1		113.6937		2 -3.63E-01	1 Unchanged Low
P51959	O12ef5	cyclin G1	116.6841	102		114.0387		2 -1.99E-01	1 Unchanged Low
Q9Y5V0		HSPC038 protein	114.0123	179	123.7928			6.54E-01	1 Unchanged Low
P56270	112ef1	MYC-associated zinc finger prote		140				-8.26E-01	1 Unchanged Low
Q9HD23		MRS2-like; magnesium homeost		117	123,8621			1 -2.72E-01	1 Unchanged Low
P78358	O03ab6	cancer/testis antigen 1	217.9167	137	123.9307			1 -6.68E-01	1 Unchanged Low
O00327	D24ab2	aryl hydrocarbon receptor nuclea		185	123.9698				1 Unchanged Low
Q9H969	F08gh5	hypothetical protein FLJ12969	89.23821	177	124.0248				1 Unchanged Low
Q15504	F01cd7	nucleoporin-like protein 1	200.4449	124				1 -6.92E-01	1 Unchanged Low
P35250	124ef6	replication factor C (activator 1)		112					1 Unchanged Low
Q9BXC0		G protein-coupled receptor 81	104,4833	128	124.3568				1 Unchanged Low 1 Unchanged Low
015232	A19ef1	matrilin 3	113.1634	110				1 -4.27E-02	1 Unchanged Low 1 Unchanged Low
076049	L19cd5	spectrin SH3 domain blinding pro		299	124.5037			1 8.00E-01	1 Unchanged Low
Q14145	C21cd8	Kelch-like ECH-associated prote		120	124,5928			1 -6.90E-01 1 7.23E-01	1 Unchanged Low
P20848	A10cd1	serine (or cysteine) proteinase in		161	124.6004				1 Unchanged Low
Q9P0N6		hypothetical protein HSPC242	112.0065	109				1 -3.53E-02 1 8.94E-01	1 Unchanged Low
P05188	E12ab2	alkaline phosphalase; placental		164					1 Unchanged Low
Q9UBJ4		transposon-derived Buster1 tran		97.3 109				2 -4.30E-01 1 -4.06E-01	1 Unchanged Low
Q13876		quiescin Q6	144,3305					1 6.78E-02	1 Unchanged Low
Q9UBF2		coatomer protein complex; subu growth differentiation factor 5 (co		115 152					1 Unchanged Low
P43026	123cd3	CGI-152 protein	111.6731	134	124.930				1 Unchanged Low
Q9HD20		RAB2; member RAS oncogene (106				2 -1.80E-01	1 Unchanged Low
Q96KK6			124.1402			3 130.7449		2 2.04E-01	1 Unchanged Low
P40617	D18ab2		127,1702	143	.23.007	, 44;	1.502-0		

PCT/US03/16096

07565·	4 140-6									
			131.989	128	125,0667	7 128,2583	-7.77E-02	-4 74E-02		1 bahanas d b
Q9Y2S			135,6724	134			-1.17E-01	2 100 02		Unchanged Low
Q1333(metastasts associated 1	104 9962	118						
Q9Y38	9 L22gh/	aminoadipate-semialdehyde de	hr 115 2252					1.71E-01		Unchanged Low
O50513	3 103gh1	UDP-Gal:belaGicNAc bela 1;4-	11 113,3332	104				-1.55E-01	1	Unchanged Low
000212		Fig homeles sees to "	5 113,705	106			1.40E-01	-1.00E-01		Unchanged Low
QBNXE			€ 407.701B	492	125,2682	2 341.5178	-1,70E+00	2.70F-01	1	Unchanged Low
			C 96.25484	146	125,3861	122,4125	3.81E-01	5 97E-01		Unchanged LOW
O95405			ta 113,9303	223			1.38E-01			Unchanged Low
P01112	K12ab	v-Ha-ras Harvey rat sarcoma vii	= 129 7328	104					1	Unchanged Low
O15120	F11cd6	1-acylglycerol-3-phosphate 0-a	0 264 2050				-4.91E-02	-3.23E-01	1	Unchanged Low
P12645	N10ab3	hone morphogenetic number 2 /	- 407.0519	238			-1,08E+00	-1.49E-01	1	Unchanged Low
P50579				117	125,4244	116.6372	2.20E-01	1.18E-01		Unchanged Low
		methlonyl aminopeptidase 2	144.5853	95.1	125.4613	121,7158	-2.05E-01	-6 04F-01	4	Unchanged Low
014908		and the protect agreeming	204,527	200			-7.04E-01	3 265 02		
QSHCC	6 117gh4	bHLH factor Hes4	82.07855	213					1	
Q15737	E23gh1	suppressor of Ty 6 homolog (S.	¢ 112 2771						1	
Q9Y519	A17cd8	chromosome 22 open reading fr	- 400 4007	140			1.62E-01	3.14E-01	1	Unchanged Low
Q9Y5J0			2 100,403/	131			3.24E-01	3.87E-01	1	Unchanged Low
P51687		mitochondrial ribosomal protein		107	125.8555	116.7986	9.99E-02	-1.33F-01	i	Unchanged Low
	O16cd1		105.2314	128	125.9497	119,5637	2.59E-01		i	Unchanged LOW
P51178	A13ef6	phospholipase C; delta 1	126,4221	162			-4.43E-03			The raining of Colo
Q9H8J6		NDRG family member 3	121.7115	198	126,048					Unchanged Low
O95900	B20ef8	hypothetical protein CLONE249	2 165 6404				5.05E-02		1	
Q9Y6N3	B01cd5	chloride channel; calcium activa	4000000	257	126.0637		-3.94E-01	6.31E-01	1	Unchanged Low
Q07869	C21cd1	porovisama antiferativa	P 99.89597	135		120,3564	3.37E-01	4.35E-01	1	Unchanged Low
Q9Y333		peroxisome proliferative activate	120,1031	112	126,2127	119.4872	7.16E-02	-9.89F-02		Unchanged Low
		chromosome 6 open reading fram	r 109.0761	114	126,2563	116,4501	2.11E-01	6.39E-02		
Q9NPD1		G protein-coupled receptor 85	206.0578	285	126,2683			4.67E-01		Unchanged Low
Q15928	K06cd3	zinc finger protein 141 (clone ph	101.1559	188	126.2732					Unchanged Low
O43826	104ab4	glucose-6-phosphatase; transpo	229 7084	111			3.20E-01	8.98E-01		Unchanged Low
Q9NWL0	305gh2	hypothetical protein FLJ20758			126.3113				1	Unchanged Low
P31629	B04ef6	himan imminadas-i	114.0453	120	126.3891			7.25E-02		Unchanged Low
O15530	110ab8	human immunodeficiency virus h	116,0174	211	126,4284	151.1911		8.64E-01	1	Unchanged Low
		3-phosphoinositide dependent pr	186.1234	193	126.45		-5.58E-01	5.15E-02	i	Linebanged Low
Q9Y2R5		milochondrial ribosomal protein !	106,3741	136	126,4713	122.8701	2.50E-01	3.52E-01		Unchanged Low
Q15208	D01cd7	senne/threonine kinase 38	158,8765	204	126,5253					Unchanged Low
Q9UET6	J19gh6	FtsJ hamolog 1 (E. coli)	102,5266	120				3.64E-01	1	Unchanged Low
Q9H5N0	F15gh5	chromosome 7 open reading fran	102 2240		126,5378			2.30E-01	1	Unchanged Low
Q9H019	B22gh4	hypothetical protein RP1-317E23	102.3345	129	126.5687	119.1911	3.07E-01	3.30E-01	1	Unchanged Low
Q9Y673	F05cd8	Aloes Commission to 1-31/E23		150	126,5974	126.5367	3.04E-01	5.53E-01	1	Unchanged Low
075690	1400ab7	Alg5; S. cerevisiae; homolog of	100.2543	131	126.6884	119,1797		3.81E-01	1	Unchanged Low
	M08gh7	UHS KerB	121.3598	150	126.7819	132.5659		3.01E-01		
Q9NUP6		hypothelical protein FLJ11220	95.908	144		122.1684				Unchanged Low
Q14296	C03cd7	FAST kinase	122.4924	92.9	126,9021			5.84E-01		Unchanged Low
Q9NW55	B20gh2	hypothetical protein FLJ10307	94.73914				5.10E-02		1 1	Unchanged Low
043639	H01cd3	NCK adaptor protein 2		141	126.9877	120.8058	4.23E-01	5.70E-01	1	Unchanged Low
P17544	L08ab2	activating transcription factor 7	139,9609	79.6	127.0293	115.5457	-1.40E-01 ·	-8.13E-01	1	Unchanged Low
075911	M01cd5	chart shall danscription ractor /	108.5839	112	127.0848	115.9726	2.27E-01	4.79E-02	1.1	Unchanged Low
Q06430		short-chain dehydrogenase/redu	156,3658	173	127.1125			1.43E-01		
	K15ef3	glucosaminyl (N-acetyl) transfera	94.56995	139	127,1541		4.27E-01			Unchanged Low
Q13769	D10cd3	chromosome 22 open reading fra	122,7812	121	127,1547		5.05E-02		1.1	Unchanged Low
Q9H864	J14gh5	hypothetical protein FLJ11021 si	97.58044		127.1909				1 1	Unchanged Low
O95500	F06cd7	daudin 14	183.0294			122.387	3.82E-01	5.45E-U1	1 1	Unchanged Low
P14222	P02cd2	perforin 1 (pare forming protein)			127.2815	163.6478 -			1 1	Unchanged Low
Q9Y4C8	B06ef2	KIAAO692 cone anniver	99.11893		127.3674	120.5243	3.62E-01	4.47E-01		Unchanged Low
Q9H3H5	M23ab4	KIAA0682 gene product	105.4769	146	127.3803	126.2296		4.67E-01		Unchanged Low
Q13433		dolichyl-phosphate (UDP-N-acet;	193.8179	108	127.5134	143.2265 -	6.04E-01 -		11	Unchanged Low
	G05cd8	LIV-1 protein; estrogen regulated	118 096		127.6823		1.12E-01 -		- 1	Hichanged Low
Q00059	007cd2	transcription factor A; mitochonds	111.4615		127.6629				1 (Inchanged Low
Q15119	106ab8	pyruvate dehydrogenase kinase;	97 01115					2.32E-01	1 (Inchanged Low
043791	F03cd3	speckle-type POZ protein				120.0132		4.81E-01	- 11	Inchanged Low
Q14206	P10cd5	Down syndrome critical region ge	111,349			117.7111	1.98E-01	3.49E-02	11	Inchanged Low
Q92926	N24cd1	SWINGAR TO THE CHICAL LEGION OF			127.7164	170.2393	9.21E-02 1	.14E+00	11	Inchanged Low
Q9P288	G15ef3	SWI/SNF related; matrix associar	115.147	105	127.7503	116.1179	1.50E-01 -	1.27E-01	11	Inchanged Low
		BRCA2 and CDKN1A interacting	148.9477	132	127.8623	136.3262 -	2.20E-01	1 725-01		
Q9NZU5	C16ef4	LIM and cysteine-rich domains 1	164.143B			139.9549	2.EOE-01 -	1.725-01		Inchanged Low
Q13569	O13cd2	Inymine-DNA glycosylase	83.16502			127.6936	3.005-01			Inchanged Low
P48167	D17ef6	glycine receptor; beta	109.5501					.05E+00	1 L	Inchanged Low
Q9NQR1	102gh4	PR/SET domain containing prote	103.0007 1E4.0405				2.24E-01 :	3.39E-01	1 (Inchanged Low
043405	B23ab4	Coordation feeter C hearty	134.2185		128.0258	124.9651 -	2.69 E-01 -7	7.35E-01	1 (Inchanged Low
Q9P028	N24ef2	coagulation factor C homolog; cc		121	128.1793	117.417		2.31E-01		Inchanged Low
		CCR4-NOT transcription comple:	104.0341	133	128.2089			3.59E-01	4.1	harageu tow
Q9Y6E8	P04cd7	sirtuin silent mating type informat	109.1684					5.37E-03	1 0	Inchanged Low
Q9NV65	H17gh3		92.33747					J.37E-03		Inchanged Low
Q9H0R8	M13gh7	GABA(A) receptor-associated pri	122,091	_		136,3658		.U3E+00	1 U	Inchanged Low
O95B17	P02cd4	BCI 2			120.3386	143.1503	7.20E-02 5	5.52E-01	1 U	Inchanged Low
P36956	K10cd2	Steroi regulatory element binding	107.1752	114	128,4319	116.5745 2	2.61E-01 S	9.05E-02	1 17	inchanged Low
	D05ab3			109	128.532	125.9988 -1	.26E-01 -3	3.60E-01	111	inchanged Low
		glucuronidase; bela	123.7102	114 1			.60E-02 -1		11	moreorges cow
	D12cd2	syntrophin; beta 2 (dystrophin-as	106.7241				2.70E-01 1			nchanged Low
	E18gh1		93.41288					.502-01		nchanged Low
	O12gh6		98.02235			122.2375 4				nchanged Low
	J03ab6	Intercellular adhesion molecule 4						.68E-01		nchanged Low
	N19gh4		101.85		128,8098		.39E-01 4		1 11	nchanged Low
	N21cd4	pulative methyltransferase	124.8741		28.8151	126,6152 4				nchanged Low
	H16cd4	beta-transducin repeal containing	152.2864	142 1	28.8641	141.0555 -2	41F-01 4	01F-01		
		G protein-coupled receptor 50	121.9424				.99E-02 -1			nchanged Low
	L24cd5		110.1643				~3C-UZ -1	.145-01		nchanged Low
	E06ef2	COLOR	96.22816			10.2029 2	.27E-01 -6	.U9E-03		nchanged Low
Q9UG66	P16ef7	hypothetical protein DKFZp586G	240 2207		20.92/2	121.3443 4	.22E-01 5	.29E-01		nchanged Low
	N17ef2			3/4 1	28.9795	273.7118 -1.	30E+00 2	.32E-01		nchanged Low
	E06cd3	Vanalass dans to	107.7937	141 1	29.1324	125.8408 2	.61E-01 3	.83E-01		nchanged Low
		very low density lipoprotein recet	113.1365	104 1	29.1355	115.483 1	91F-01 4	10F-01		
	C07gh5	1ERA protein	94.63638	199 1	29,2792 4	140.8474 4	SOC 04	075+00		nchanged Low
	C01ab8	peroxisomal membrane prolein 3 3	321 6292	219 1	29 3474 -	222 222	.JUE-UI 1,	U/ E*U0		nchanged Low
	M24cd6	ublquitin-conjugating enzyme E2	117 2071	100 1	20 25	223.2274 -1.	ა1⊵+00 -5	.56E-01		nchanged Low
		inter-alpha (globulin) inhibitor; H: 2	20 4455	100)	29.3512 1	115.6428 1	.42E-01 -2	.24E-01		nchanged Low
	M18ab8	Directorist and a street of the second	47.00	100 1	29.3613 1	185.0551 -A	88F-01 .3	62E-01		nchanged Low
		phosphatidylinositol 4-kinase; ca	117.2574	115 1	29.4365 1	20.4316 1	.43E-01 -3	31E-02		ichanged Low
						•			. 0	

Q9Y5J6	J16cd7	fracture callus 1 homolog (rat)	102,357	127	129.4503		3.39E-01	3.06E-01	1 Unchanged Low
Q92889	M24ab4	excision repair cross-complemen		97.6	129.5255		-2.83E-01	-6.91E-01	1 Unchanged Low
P08697	C07ef1	serine (or cysteine) proteinase in		136	129,5425			-2.29E-02	1 Unchanged Low
Q9UBX1 Q9NPI7	A05cd4 D22eft	calhepsin F	113.4065	118	129.6557			6.08E-02	1 Unchanged Low
P11586	D14ef7	hypothelical protein LOC51315 methylenetetrahydrofolate dehyd	240.5378	171 106	129.76 129,8444		-8.90E-01		1 Unchanged Low
096010	J15cd4	LIM domain binding 1	90.9506	155	129,8567		-1.21E-01 5.14E-01	-4.11E-01 7.67E-01	1 Unchanged Low
Q13188	P13cd2	serine/threonine kinase 3 (STE2)		135	129.8809		1.45E-01	1.99E-01	1 Unchanged Low 1 Unchanged Low
Q13845	121ab3	B-cell CLL/lymphoma 7B	145.9359	88.3	129,9273				1 Unchanged Low
Q9Y5W9	D07cd8	sorting nextn 11	144.6605	198	130.0152			4.56E-01	1 Unchanged Low
060539	102cd7	Ras association (RalGDS/AF-6)	126.8466	102	130.0346	119.7412		-3.10E-01	1 Unchanged Low
Q60885	102ef3	bromodomain-containing 4	114.9928	98.1	130.1093		1.78E-01	-2.29E-01	1 Unchanged Low
Q9H2B0 P51582	O16gh5	nucleosome assembly protein 1-1		147	130.1112		8.60E-02	2.61E-01	1 Unchanged Low
Q00585	C24ab8 F08cd1	pyrimidinergic receptor P2Y; G-p		109	130.158			-9.03E-02	1 Unchanged Low
Q15131	B23ef5	small inducible cytokine subfamil cyclin-dependent kinase (CDC2-		216 108	130.1849		-3.74E-01	3.53E-01	1 Unchanged Low
075148	F08cd8	cryptochrome 2 (photolyase-like)		172	130,2201		-3.73E-01 5.97E-01	9.99E-01	1 Unchanged Low
Q15691	G17cd8	microlubule-associated protein; F		168	130.2758		-4.15E-01		1 Unchanged Low
Q15779	H21ab8	procollagen (type III) N-endopepi		133	130.2905		-1.18E-01	-8.51E-02	1 Unchanged Low 1 Unchanged Low
Q9UHD2		TANK-binding kinase 1	82.31867	172	130,2947			1.06E+00	1 Unchanged Low
Q9NSS3	K15gh3	hypothetical protein DKFZp434E		115	130,3095	117.5518	2.77E-01	9.44E-02	1 Unchanged Low
Q13797	101ef7	integrin; alpha 9	156.9946	85	130.4752		-2.67E-01		1 Unchanged Low
P46531 043262	M10e/5 C01cd6	Notch homolog 1; translocation-e		164	130,4897		6.01E-01	9.30E-01	1 Unchanged Low
Q9Y2H6	K12ef8	deleted in lymphocytic leukemia; KIAA0970 protein	170.8655	195 128	130,4943 130,5238		7.65E-01		1 Unchanged Low
P25788	J11ab8	proteasome (prosome; macropali		285	130,5238		-3.89E-01 -9.86E-01	-4.22E-01 1.40E-01	1 Unchanged Low
Q9BQ\$8	D09gh7	FYVE and coiled-coil domain cor		207	130,6451	167.5571	-3.34E-01	3.33E-01	1 Unchanged Low 1 Unchanged Low
Q9Y396	J04ef1	elongation of very long chain fatt		82.2	130.6722			-9.85E-01	1 Unchanged Low
Q9P0R7	K06ef2	hypothetical protein LOC51242	126.7302	218	130,6804		4.43E-02	7.84E-01	1 Unchanged Low
015269	F17cd6	serine palmitoyltransferase; long		133	130,7429	120.3129	4.24E-01	4.46E-01	1 Unchanged Low
Q9UFX0	B02ef2	calcium binding atopy-related au		287	130.8943		-8.35E-01	2.98E-01	1 Unchanged Low
Q9H663 075410	F24gh5 P17cd2	likely ortholog of mouse actin-rela		128	130,9258		4.15E-01	3.87E-01	1 Unchanged Low
Q9Y5N5	114cd8	transforming; acidic colled-coll co putative N6-DNA-methyltransfera		133	130.9827		-7.75E-01		1 Unchanged Low
Q98Q72	F02gh7	hypothetical protein MGC10471	106.3824	133 115	131.0924 131.1126	121,2285 117,4135	4.00E-01	4.24E-01	1 Unchanged Low
Q9Y303	E10ef2	CGI-14 protein	153.5787	101	131,2035		3.02E-01 -2.27E-01	1.09E-01	1 Unchanged Low 1 Unchanged Low
Q9P005	A22ef8	HSPC159 protein	341.3083	579	131.3148		-1.38E+00	7.64E-01	1 Unchanged Low 1 Unchanged Low
Q9Y3A1	H19ef2	DKFZP566O084 protein	200.4927	262	131.514		-6.08E-01	3.88E-01	1 Unchanged Low
Q15172	C12cd1	protein phosphalase 2; regulator		140	131,5524	146.553	-3.56E-01	-2.69E-01	1 Unchanged Low
Q98V19 Q14534	119gh6	hypothetical protein MGC955	96.47316	141	131.6188		4.48E-01	5.43E-01	1 Unchanged Low
Q9BS43	E11cd2 G19gh8	squalene epoxidase hypothetical protein MGC12435	155.5799	71.3	131,8107	119.574	-2.39E-01	-1.13E+00	1 Unchanged Low
P49788	D24ab8	retinoic acid receptor responder	592.1694 97 19637	121 135	131.9022 131.9355	121,3412	-2.17E+00 4.41E-01		1 Unchanged Low
Q9UP83	M10cd6	component of oligomeric golgi co		141	131,936		3.96E-01	4.73E-01 4.89E-01	1 Unchanged Low 1 Unchanged Low
Q9Y2X0	N11cd5 ·	thyroid hormone receptor-associa		97.9	131,9541	115.5741	1.75E-01		† Unchanged Low
Q9Y3E3	115ef2	CGI-145 protein	94.76733	132	131.9919	119.5355	4.78E-01	4.76E-01	1 Unchanged Low
095183	J14cd6	veside-associated membrane pn		157	132.0539	126,6473	5.34E-01	7.81E-01	1 Unchanged Low
Q13242	N22cd3	splicing factor; arginine/serine-rk		141		143.0016	-2.41E-01	-1.48E-01	1 Unchanged Low
Q9Y5R5 P04181	A06gh1 N02ab7	doublesex and mab-3 related trai		162	132.1528	131.5571	3.90E-01	6.81E-01	1 Unchanged Low
Q14254	B15ab4	omithine aminotransferase (gyra- flotillin 2	130.1586	106 154	132,1687 132,2038	116.724		-8.99E-02	1 Unchanged Low
Q9Y6E6	P08cd7	sirtuin silent mating type informat		136	132,2501	138,6301 121,0265	2.25E-02 4.86E-01	2.38E-01 5.31E-01	1 Unchanged Low
Q9H237	D20gh5	porcupine	148.2204	258	132,2656		-1.64E-01	7.98E-01	1 Unchanged Low 1 Unchanged Low
Q92785	N21cd2	requiem; apoptosis response zin-		108	132,3893	119.7533	1.61E-01		1 Unchanged Low
095458	N11cd2	tubulin-specific chaperone d	128.9522	100	132,4761	120.4814	3.89E-02		1 Unchanged Low
P20749 Q9H2N8	P23ef6	B-cell CLL/lymphoma 3	264.1174	186	132.484		-9.95E-01		1 Unchanged Low
Q15020	D24gh7 M08gh1	hypothelical protein GL012	104.3233	160	132.5137	132.1507	3.45E-01	6.14E-01	1 Unchanged Low
P42025	O01gh1	squamous cell carcinoma antiger ARP1 actin-related protein 1 hon		147 92	132,5393	132.2727	1.77E-01	3.26E-01	1 Unchanged Low
P98173	123gh5	2.19 gene	113.0426	116	132,554 132,6703	156.1129 120.727	-8.79E-01 2.31E-01	4.31E-02	1 Unchanged Low 1 Unchanged Low
095295	C06cd8	SNARE associated protein snapt		119	132,6986		-1.29E+00		1 Unchanged Low
Q9NVA1	F15gh3	chromosome 20 open reading fra		100		114.8529	2,48E-01		1 Unchanged Low
P35573	O03ab2	amylo-1; 6-glucosidase; 4-alpha-		106		116.8065			1 Unchanged Low
Q13126	K13ab7	methylthioadenosine phosphoryli		236		190.4847		2.17E-01	1 Unchanged Low
P43034 Q01844	C11ab8 E15ab5	platelet-activating factor acetylhy		125		121,4511		2.22E-01	1 Unchanged Low
P27482	E18ab6	Ewing sarcoma breakpoint region calmodulin-like 3		102	133,001	115.5021	2.57E-01		1 Unchanged Low
Q9NUW4		BRIX	114.7017 223.8591	104 77.1		117.2725	2.14E-01		1 Unchanged Low
P78552	J23ab6	interleukin 13 receptor; alpha 1	118.2476	93.9	133.0208 133.0765	115,0749	-7.51E-01 1.70E-01		1 Unchanged Low
P51843	O24ab2	nuclear receptor subfamily 0; gro			133,1401		3.46E-01		1 Unchanged Low 1 Unchanged Low
P15822	L23ab5	human immunodeficiency virus h				125.3978	6.78E-01		1 Unchanged Low
Q9BXY9	O13ghB	RALBP1 associated Eps domain		140	133,274	121,2561	5.53E-01		1 Unchanged Low
Q9ULB4	P10ef1	cadherin 9; lype 2 (T1-cadherin)		137	133,4641		-1.04E+00 ·	-1.01E+00	1 Unchanged Low
Q9NZZ7 Q02086	C08ef8 C09cd2	HSPC171 protein	214.9523		133.5412			3.04E-01	1 Unchanged Low
Q98UV9	H14gh6	Sp2 transcription factor DKFZP586J0119 protein	101.0989	133		122.3826	4.02E-01	3,90E-01	1 Unchanged Low
Q9UHJ9	A09ef4	FGF receptor activating protein 1	102.1174	119 121	133,5923		3.88E-01	2.22E-01	1 Unchanged Low
Q16401	E11cd1	proleasome (prosome; macropali			133,6008	132,2946	4.07E-01		1 Unchanged Low 1 Unchanged Low
Q9Y4K4	D04cd6	mitogen-activated protein kinase			133.6676		1.37E-02		1 Unchanged Low
O60463	H06cd3	phosphatidic acid phosphatase to	103.2417			129.2479	3.73E-01		1 Unchanged Low
095639	P20cd6	cleavage and polyadenylation sp		67.6	133.7241	128,1539	-4.54E-01 -	1.44E+00	1 Unchanged Low
P50539	N19ab7	MAX interacting protein 1	175.6646	187	133.7304	165.4016	-3.93E-01	8.87E-02	1 Unchanged Low
Q9UH62 Q15525	115ef3 C13ab7	ALEX3 protein	118.2527		133.7314		1.77E-01	5.32E-01	1 Unchanged Low
Q9H8T0	H21gh5	v-maf musculoaponeurotic fibros: fused toes homolog (mouse)	105.3252		133,7521		3.45E-01		1 Unchanged Low
Q9Y4X5	K03cd6	ariadne homolog; ubiquitin-conju			133,7606	160.2791 120.4976	3.33E-01		1 Unchanged Low
					.00,011	140,4310	7.20E-U1	J.00E-01	1 Unchanged Low

Q9H9E1	O01gh5	ankyrin repeat; family A (RFXA)	N 172,6266	78.	R 134.053	2 128.4841	1 -3 65E 0	1 125,00			Heat	
Q93063	F13ab3	exostoses (multiple) 2	123.2886	16	1 134.3466	139.4501		3.82E-01			Unchange Unchange	
Q9H992			89,54768	14				7.37E-01			Unchange	
043237	P02ab4		₩ 290.6769	111				-1.28E+00		i	Unchange	ed Low
Q99643 P24386	H02cd1 F11ef1		di 143.4474	12	5 134.5527			-1.95E-01		1	Unchange	d Low
Q9Y5F7	F12gh4	chorolderemia (Rab escort prot	el 93,86058	12:				4.56E-01			Unchange	
P06756	D14ab6		/ 101.7878	11-							Unchange	
Q9H5L7	C02gh6		£ 200.0782	219			-5.72E-01				Unchange	
015517	A14ef6	turnor necrosis factor receptor s	173.0100	140 201				-2.34E-01			Unchange	
Q9P0N5		HSPC244	107.9991	129				1.17E+00			Unchange	
Q9NWT2		hypothetical protein FLJ20623	107.8864	105				2.53E-01 -4.05E-02			Unchange	
Q9Y2A0	002cd7	TP53 target gene 1	127,408	88.2				-5.30E-01			Unchange	
Q9NZ43	N16gh4		si 141.2528	137				-4.24E-02			Unchange Unchange	
P27986	A10ef1	phospholnositide-3-kinase; regu		112							Unchange	
Q9UBP6 Q9BWK5		methyltransferase-like 1	107.2187	122				1.85E-01		i	Unchange	d Low
P54802	M24ab6	hypothetical protein MGC5242	101.1324	123			4.175-01	2.88E-01			Unchange	
075963	H09cd7	N-acetylglucosaminidase; alpha G-protein coupled receptor		66.3				-1.64E+00		1	Unchange	d Low
Q9NXZ4	F03ef4	ELG protein	110.3886 124.5205	184 105				7.40E-01			Unchange	
060353	B11cd3	frizzled homolog 6 (Drosophila)	85.64918	146			-	-2.50E-01			Unchange	
Q9NQS5		latexin protein	94,21822	163				7.73E-01 7.90E-01			Unchange	
014776	A19cd7	transcription elongation regulate	113.3803	117						1	Unchange Unchange	d Low
Q9Y5A2	L13ef1	pulative zinc finger protein NY-F	154.0704	188			-1.83E-01			11	Unchange	d Low
P49842 Q06546	G23cd6	serine/threonine kinase 19	133.3557	171	135.7013			3.56E-01			Unchange	
Q9Y2E7	J14ab5 102ef8	GA binding protein transcription		138			6.10E-01	6.34E-01			Inchange	
Q9UM11	102ef2	KIAA0938 protein Fzr1 protein	109.8218	105				-7.00E-02			Inchange	
Q99447	B14ab8	phosphate cytidylyltransferase 2	80.82847	186			7.50E-01			1 (Jnchange	d Low
Q9H1R3	F22gh8	myosin light chain kinase 2; skel	, 174.0 <u>2</u> 01	77.4 129				-1.18E+00			Inchange	
P28358	P05ab6	homeo box D10	108.9905	102				4.63E-01 -9.71E-02			Inchange	
076070	P18cd1	synuclein; garnma (breast cance	128,551	140				1.18E-01			Inchange	
Q9Y5A7	N07ef2	NEDD8 ultimate buster-1	119.0653	195		149.951		7.08E-01			Jnchanger Jnchanger	
Q9NWY7		hypothetical protein FLJ20530	103.7638	110	136.3026			8.20E-02			Inchange	
043715 Q9BY75	P02ef2	hypothetical protein HSPC132	137.4694	110		127.9562	-1.23E-02			1 l	Inchange	tlow
043431	K19gh8 A12e/7	itchy homolog E3 ubiquitin protei	118.5721	127		127.3323	2.03E-01				Inchange	
043567	D15cd7	endothelial differentiation; lysopt ring finger protein 13		171			1.116-01	4.34E-01			Inchanged	
Q9Y2S7	P13ef8	DKFZP586F1524 protein	110.4449 113.1366	153 104		133.3964	3.06E-01	4.72E-01			Inchanged	
014582	O24ef2	spondyloepiphyseal dysplasia; la	135,1666	135		117.8414 135.7349		-1.26E-01			Inchanged	
075056	G12gh1	syndecan 3 (N-syndecan)	120.0661	162		139,562	1.66E-02 1.88E-01	1.46E-03 4.31E-01			Inchanged	
Q9P2Y2	D18gh4	beta-1;4 mannosyltransferase	89.6857	143		123.2251	6.09E-01				Inchanged Inchanged	
Q16595	M08ab3	Friedreich ataxia	139.5657	130		135.5917	-2.85E-02	-9.82E-02			Inchanged	
Q14807 Q9BZL4	B18ab7	kinesin-like 4	114.4726	94.2	136.9298	115.2048				iù	Inchanged	Low
P30876	L10gh6 I13ab8	protein phosphalase 1; regulator	110.5367	123		123,5234		1.54E-01	1	ı	Inchanged	Low
Q9UGP6	B17cd7	polymerase (RNA) II (DNA direct coatomer protein complex; subur	265.5337	200		200.7168			1		Inchanged	
P52655	M01ef2	general transcription factor ItA; 1	121 / 200	134 128	137.3091	122.0845	5.35E-01	5.02E-01		l	Inchanged	Low
Q9HB07	G23gh5	chromosome 12 open reading fra	121.4650	118	137.3707 137.4723	129.1154	1.77E-01	8.08E-02	1		Inchanged	
Q92940	H23ab7	MAD; mothers against decapents	414.4889	223	137.6028	125.5248 258.3032		-4.11E-02	1		Inchanged	
Q9NWN1		hypothetical protein FLJ20727	117.7514	120	137,6351		2.25E-01		1	1 1	Inchanged Inchanged	LOW
O43664 O14647	A10cd6	G protein-coupled receptor 66	108.1311	102	137.6789	116,0604	3.49E-01		1	i	nchanged	Low
075333	C19ab4 N15cd2	chromodomain helicase DNA bin T-box 10		164		132,5407	5.19E-01		i	U	nchanged	Low
Q16539	J04ef5	mitogen-activated protein kinase	126.0065	129	137.7737	130,9257	1.29E-01	3.38E-02	1	U	nchanged	Low
	A07gh5	chromosome 20 open reading fra	86 80552	219 143	137.8	168.4312		5.57E-01	1	U	nchanged	Low
Q15149	E05ab8	plectin 1; Intermediate filament bi	125.7248	83.9	137,803 138,0785	122.4598 115.8848	6.67E-01	7.18E-01	1		nchanged	
Q9NVZ7	G12gh3	hypothetical protein FLJ10407	166.213	175	138.1444	159.6342	1.35E-01		1		nchanged	
	D02cd2	steral-C4-methyl oxidase-like	171.8648	71.2	138.1497	127.0641		7.06E-02	1		nchanged	
	D16gh5	hypothetical protein FLJ12387 si	96.54943	129		121.3572	5.17E-01	4.22E-01			nchanged nchanged	
	K14cd7 D24cd3	mitochondrial ribosomal protein t	94.17331	128		120.1502			i		nchanged	
	J19ef7	KH-type splicing regulatory prote	126.0509	88.3	138.3014	117.5428	1.34E-01	-5.14E-01	1	Ū	nchanged	Low
	B14cd2	aminopeptidase puromycin sensi sodium channel; voltage-gated; t	273.7328	98.9	138.3924	170.346	-9.84E-01	1.47E+00	1	Ų	nchanged	Low
	M01ef7	erythropoietin receptor	198.5954	139	138.4331	134.9928	1.20E-01	1.28E-01			nchanged	
015228	A16ef3	glyceronephosphate O-acyltrans		77.3 186	138.4803 138.5137	138.1099	-5.20E-01 -	1.36E+00	1		nchanged	
	K05cd8	F-box only protein 9	151.0501	104	138,5593	183.349	-7.01E-01 -1.25E-01			U	nchanged	Low
	D20cd6	chaperonin containing TCP1; sut	274.982	178		197.1593					nchanged	
	P19cd8	fer-1-like 3; myoferlin (C. elegans	218,8411			186.4122					nchanged nchanged	
	N20cd1	SWI/SNF related; matrix associa	131.7669	82.7	138.6764	117.704	7.37E-02		i	U	nchanged	low
	A18gh1 10gh4	sterol-C5-desaturase (ERG3 delt			138.7232	131.1196	4.45E-01				nchanged	
	C19ab2	chromosome 12 open reading fra amiloride-sensitive cation channe	80.2401	158	138.8274	125.605	7.91E-01		1	U	nchanged	Low
	F16ab5	CDC-like kinase 1	86.24615		138.9915			9.06E-02	1	Ų	nchanged	Low
P33552	A22ab5	CDC28 protein kinase 2	109.5671	147 96	139.0985		6.90E-01		1	U	nchanged	Low
O75808	H21cd2	small optic lobes homolog (Drosc	115,654	98.9		114.8881	3.45E-01		1	u	nchanged	Low
	M07gh2	LIS1-interacting protein NUDE1:	96.20481		139.1939	117.8898 119.5057	4.07 E-U1	2.20E-U1	1	U	nchanged	Low
	L12ab8	peroxisomal biogenesis factor 14	473.8395	113	139.2209	241.8793 -	1.77E+00 -	2 07E+00	- 1	U	nchanged nchanged	LOW
	P23cd1	ribonuclease H1	167.3611	111	139.2393	139.1464	-2.65E-01	5.95E-01			nchanged i	
	P14cd4	golgi SNAP receptor complex me	115.7917	89.7	139,2541	114.9152	2.66E-01	3.68E-01			ichanged i	
	L24gh1 P06cd2	milochondrial ribosomal protein L		120	139,3426	156.1365 -	-5.87E-01	8.04E-01			changed	
		ras-related C3 botulinum toxin su	158.176	166	139,3634	154.6374	-1.83E-01	7.29E-02			changed	
	109ef2	tubulin-specific chaperone c CGI-127 protein	103.3699 B3.4762				4.34E-01		1	U	changed	Law
P18827	F22cd1		114,4045		139.6599	128.8568	7.42E-01	9.69E-01	1	Ur	changed I	Low
	321ab3	nuclear receptor subfamily 3; gro	106.6921	92.7 103	139,6607	110.0895	2.88E-01	3.03E-01	1	Ų	changed	Low
		, -, 9,0		103	139.6889	110.0017	3.03E-01 ·	4,495-02	1	Uſ	ichanged I	Low

Q9Y301	N05ef1	CGI-12 protein	108.0511	101	139.7035	116.3594	3.71E-01	-9.27E-02	1	Unchanged Low
000626	F10cd1	small inducible cytokine subfamil	113.478	106	139.8189	119.8588	3.01E-01	-9.45E-02		Unchanged Low
	B1Bcd8	neural precursor cell expressed;	108.1711	133	139,9597	127.048	3.72E-01		1	Unchanged Low
Q9Y311 .	C01ef2	mitochondrial ribosomal protein t	162.8519	74.8	139,9856		-2.18E-01			Unchanged Low
Q15633	K14cd2	TAR (HIV) RNA binding protein 2		125	140.0138		-2.75E-01			Unchanged Low
P02278 P53672	M22gh6 J11cd7	H2B histone family; member K crystallin; beta A2	100.2794 101.7321	216 187	140,0965 140,2804	152.2309 142,8494	4.82E-01			Unchanged Low
P09417	C05ab8	quinoid dihydropteridine reductas		161	140.2664	142.5395	4.64E-01 1.48E-01	8.75E-01 3.43E-01		Unchanged Low
Q9HAY2	K14gh5	MAGEF1 protein	96.01675	126	140.3654	120.9341	5.48E-01	3.97E-01		Unchanged Low Unchanged Low
P30273	A20ab5	Fc fragment of IgE; high affinity I		171	140,4541	133.3475	6.67E-01			Unchanged Low
Q92949	G10ab4	forkhead box J1	250.4112	167	140.5084	185,89	-8.34E-01			Unchanged Low
Q9GZT9	G01gh7	egl nine homolog 1 (C. elegans)	112.8025	99.2	140.6556	117.5626	3.18E-01	-1.85E-01		Unchanged Low
P35658	O23cd4	nucleoporin 214kD (CAIN)	141.8279	137	140.7761	139.8811	-1.07E-02	-4,96E-02	1	Unchanged Low
Q9HB40	G01gh5	likely homolog of rat and mouse :		99.3	140.9548	115.3964		-9.32E-02		Unchanged Low
014686	O16cd3	myeloid/lymphoid or mixed-linear	117.844	157	141.0674	138.5398		4.11E-01		Unchanged Low
Q92911 Q9Y5E4	O12cd1 B18ef3	solute carrier family 5 (sodium to- protocadherin beta 5	95.08061	125 148	141.0904 141.1894	176.594 127.9379	-9.05E-01 5.70E-01	-1.08E+00		Unchanged Low
Q13057	H17gh7	nucleotide binding protein	186.8796	125	141.2202		-4.04E-01		1	Unchanged Low Unchanged Low
Q14863	B23ab8	POU domain; class 6; transcriptic		103	141.3258	115.7954		-4,97E-03		Unchanged Low
P36610	G10ef3	frequenin homolog (Drosophila)	110.2191	136	141.4925	129.0874	3.60E-01	2.98E-01		Unchanged Low
060240	O12ab8	perilipin	1121.108	52.2	141.5046	438.2653	-2.99E+00	-4.43E+00		Unchanged Low
Q9H9B4	P05gh5	hypothetical protein FLJ12876	87.99841	133	141.7125	120.8973	6.87E-01	5.96E-01	1	Unchanged Low
000584	J14cd3	ribonuclease 6 precursor	124.6932	250	141.8008	172.0542		1.00E+00		Unchanged Low
Q9H658 P18858	M04gh7 K14ef6	G protein-coupled receptor 107	158.2595	214	141.8139		-1.58E-01	4.32E-01		Unchanged Low
Q02877	D23cd1	ligase I; DNA; ATP-dependent ribosomal protein L26	92.33071 449.1728	128 297	141.8761 141.8851	120.8853	6.20E-01	4.76E-01 -5.99E-01		Unchanged Low
P15586	J07ab4	glucosamine (N-acetyl)-6-sulfata:	261.466	119	141.968	174.2861		-1.13E+00		Unchanged Low Unchanged Low
P42696	H08ef4	KIAA0117 protein	110.9202	231	141.9948	161.4649		1.06E+00		Unchanged Low
075710	A17ef1	molybdenum cofactor synthesis *	105.3087	101	142,0352	116.2449		-5.47E-02		Unchanged Low
Q9NPI8	C01gh2	Fanconi anemia; complementatic	193.8104	214	142.0643	183.3737	-4.48E-01	1,45E-01		Unchanged Low
Q9HAR6	O10gh5	uterine-derived 14 kDa protein	118.9551	198	142.0669	152.8734	2.56E-01	7.32E-01		Unchanged Low
P15018	O18ef7	leukemia inhibitory factor (cholini		309	142.2733		-8.35E-01	2.84E-01		Unchanged Low
Q13546 P24298	A23cd4 H01ab4	receptor (TNFRSF)-interacting suglutamic-pyruvate transaminase •		131	142.2746			-9.15E-02		Unchanged Low
Q14129	C22cd4	DiGeorge syndrome critical regio	210.426	97.† 141	142.3123 142.4302	149,9584 131,8973	3.45E-01	-1.12E+00 3.32E-01		Unchanged Low
Q9BQA2	K09gh8	KIAA1882 protein	164.9357	187	142,4325		-2.12E-01	1.80E-01		I Unchanged Low I Unchanged Low
Q9UJX9	B12ef3	hypothetical protein; estradiol-inc		98.5	142.5053			-3.06E-01		Unchanged Low
Q99590	101cd5	splicing factor; arginine/serine-ric		199	142,5273			-2.18E-01		Unchanged Low
O14798	C06ef7	tumor necrosis factor receptor su	142.6951	213	142.5294	166.073	-1.68E-03	5.78E-01		Unchanged Low
P32238	P01ab3	cholecystokinin A receptor	487,368	168	142,5911			-1.54E+00		Unchanged Low
Q9UQ90	C17cd2	spastic paraplegia 7; paraplegin	101.2857	146	142.6321	129.9052		5.26E-01		Unchanged Low
P82980 P49716	M03gh8 P15ab5	retinol binding protein 5; cellular CCAAT/enhancer binding proteir	111.2191	93.9 73.4	142.6923	115.9454		-2.44E-01		Unchanged Low
P20062	M12cd1	transcobalamin II; macrocytic and	125.6347	141	142.8277 142.8402	128.4871 136.6279	1.85E-01	-1.20E+00 1.71E-01		I Unchanged Low I Unchanged Low
P21580	B05cd2	turnor necrosis factor; alpha-indu		317	142.8664		-9.49E-01	1.99E-01		Unchanged Low
Q99774	H01cd5	protein phosphatase 4; regulator	195.85	173	143.1867			-1.76E-01		Unchanged Low
Q9P0T1	A17e/3	hypothetical protein HSPC192	104.4363	101	143.2517	116.1881	4.56E-01	-5.00E-02		1 Unchanged Low
P06737	B18ab8			139	143.2732	122.6343	7.51E-01			Unchanged Low
015063	K10gh1	KIAA0355 gene product	149.9191	144	143.2945			-5.91E-02		1 Unchanged Low
P02743 Q9NVT0	K14ab2 I14gh3	amyloid P component; serum hypothetical protein FLJ10534	108.0152 136.4264	94.7 109	143.3031 143.3056	115.3343 129.6057		-1.90E-01		Unchanged Low
Q9Y4P3	O07ef3	transducin (bela)-like 2	112.0852	104	143.4853	119.7013		-3.23E-01 -1.14E-01		1 Unchanged Low 1 Unchanged Low
Q13886	N14ab3	basic transcription element bindit	164.4049	88.7	143.5269			-8.90E-01		1 Unchanged Low
Q9NVP8	L19gh4	hypothetical protein FLJ10595	101.7956	108	143,5829		4.96E-01			1 Unchanged Low
P08240	G05cd2	signal recognition particle recept-	158.3565	132	143.6823	144.5673	-1.40E-01	-2.66E-01		1 Unchanged Low
P49459	M19ef5	ubiquitin-conjugating enzyme E2.	98.6462	170	143.74	137.6267	5.43E-01	7.89E-01		1 Unchanged Low
Q9NVA8	L16gh2	solute carrier family 38; member	112.6488	127	143.7595	127.9498	3.52E-01	1.78E-01		t Unchanged Low
P25325	K23cd3 K08cd7	mercaptopyruvate sulfurtransfera		274	143.8077			-7.45E-02		1 Unchanged Low
Q9UHI6 P52298	H01cd7	DEAD/H (Asp-Glu-Ala-Asp/His) t nuclear cap binding protein subu	138.9223	209 191	143.8383 143.8819	162.7208 157.996	8.43E-02 5.06E-02			f Unchanged Low
Q9NQ30	A10cd7	endothelial cell-specific molecule	104.1846	122	143,9703	123.2921	4.67E-01	2.24E-01		1 Unchanged Low f Unchanged Low
O00541	108ef3	pescadillo homolog 1; containing	180.43	204	144.0988			1.78E-01		1 Unchanged Low
Q9H2V9	B06gh7	hypothetical protein CDA08	173.5608	169	144.1057			-3.56E-02		1 Unchanged Low
Q9NNX1	F12ght	tuftelin 1	147.0323	175	144.106	155.2699	-2.90E-02	2.49E-01		1 Unchanged Low
Q13508	H14ab2	ADP-ribosyltransferase 3	109.4052	101	144.1545	118.095	3.98E-01	-1.19E-01		1 Unchanged Low
Q9NR30	B21gh6	DEAD/H (Asp-Glu-Ala-Asp/His) t		130	144.2348	128.2929		2.36E-01		1 Unchanged Low
Q14320	E15cd5 D03cd2	DNA segment on chromosome X		178	144.2677	141.6929		7.99E-01		1 Unchanged Low
Q04726 P20061	N13cd1	transducin-like enhancer of split: transcobalamin I (vitamin B12 bir		115 85.5	144.3217 144.4835	156.9544 115.787		-8.86E-01 -4.56E-01		1 Unchanged Low
Q9Y2Y1	J12ef2	polymerase (RNA) III (DNA direc		145	144.5561			-8.58E-02		1 Unchanged Low 1 Unchanged Low
P35754	G22ab5	glutaredoxin (thioltransferase)	153.823	138	144.7308			-1.59E-01		1 Unchanged Low
Q02978	F01cd3	solute carrier family 25 (mitochor	106.877	98.2	144,7822	116.628		-1.22E-01		1 Unchanged Low
095792	N20gh2	protein associated with PRK1	130.6914	146	144.7909	140.5872	1.48E-01	1.63E-01		1 Unchanged Low
Q13232	D20ef7	non-metastatic cells 3; protein ex	90.7906	156	144.8861	130.592		7.82E-01		1 Unchanged Low
Q9NVH2	L01ef8	DKFZP434B168 protein	117.6587	91.2	144.9821	117.932		-3.68E-01		1 Unchanged Low
Q15561 Q9UPN9	O15cd2 B01ef2	TEA domain family member 4 tripartite motif-containing 33	234,306 110,9269	172	145.3307			-4.46E-01		1 Unchanged Low
Q90PN9 Q14353	006ab3	guanidinoacelate N-methyltransfi		135 103	145.3321 145.3798	130.2668 119.1987		2.78E-01 -8.24E-02		1 Unchanged Low 1 Unchanged Low
Q9NZM5	A18ef4	glioma turnor suppressor candida	150.098	122	145.587			-3.01E-01		1 Unchanged Low 1 Unchanged Low
043255	B11cd2	seven in absentia homolog 2 (Dn		101	145.653			-1.81E-01		1 Unchanged Low
Q14331	G04ab5	FSHD region gene 1	164.0964	176	145,6576		-1.72E-01			1 Unchanged Low
P18847	N23ab5	activating transcription factor 3	127.5442	154	145.6654	142.5299		2.75E-01		1 Unchanged Low
O60258	107cd4	fibroblast growth factor 17	159.7671	223	145.6821		-1.33E-01	4.79E-01		1 Unchanged Low
P47929	P02ab6	lectin; galacloside-binding; solub		92.8	145.699			-7.04E-01		1 Unchanged Low
P35227 Q15599	O14cd4 O15cd5	zinc finger protein 144 (Mel-18)	108.2295	151	145.8554			4.80E-01		1 Unchanged Low
Ø 10033	J	solute carrier family 9 (sodium/h)	100.30/3	80.5	145.993	131.8082	-2.11E-01	-1.07E+00		1 Unchanged Low

Q9UKJ5	B20cd7	cystein-rich hydrophobic domain		206	145,9967	178.8637	-3.37E-01	1.60E-01	1 Unchanged Low
Q9HCB9	C10gh5	chromosome 1 open reading fran	94.16759	126	146,1683	122.1779	6.34E-01	4,22E-01	1 Unchanged Low
Q9Y3A0	K07ef2	CGI-92 protein	119.1033	178	146.1698	147.7783	2.95E-01	5.80E-01	1 Unchanged Low
P49815 P25116	M17ef6	tuberous sclerosis 2	106.0159	163		138,4231	4.64E-01	6.20E-01	1 Unchanged Low
075829	L19ab4 O01cd7	coagulation factor II (thrombin) re		143	146.2742	123.8585	8.27E-01	7.92E-01	1 Unchanged Low
Q9NYI1	M16ef2	chondromodulin I precursor ECSIT	121.7658 111.7926	119 110	146.4876	128,9495	2.67E-01		
P24347	124ef5	matrix metalloproteinase 11 (stro		136	146.6345 146.649	122.8588 126.9018	5.88E-01	-2.14E-02 4.84E-01	1 Unchanged Low 1 Unchanged Low
043189	K24ab8	PHD finger protein 1	112.1101	101	146.6925	120,038		-1.46E-01	1 Unchanged Low
Q9P029	N22ef2	TH1-like (Drosophila)	201,2292	159	146.6998		-4,56E-01		1 Unchanged Low
Q9NZP9	E17ef2	immediate early response 5	105.1683	157	146.761	136,3041	4.81E-01	5.78E-01	1 Unchanged Low
Q9UIA3	112e/2	neutral sphingomyelinase	165.6408	147	146.8134	153.0756	-1.74E-01		1 Unchanged Low
Q99734	M12ef5	Notch homolog 2 (Drosophila)	220.4251	263	147.0111	210.2011	-5.84E-01	2.56E-01	1 Unchanged Low
O00411 P28347	C19cd1	polymerase (RNA) mitochondrial	120.5078	105	147.0481	124.1084		-2.02E-01	1 Unchanged Low
Q13619	P04gh1 G19gh1	TEA domain family member 1 (S) cullin 4A		174	147.0913	183.776	-6.46E-01		1 Unchanged Low
094929	O06ef8	KIAA0843 protein	153.1592 108.7821	218 99.4	147,2921	172.6621		-1.31E-01	1 Unchanged Low
Q9P026	A02ef8	HSPC134 protein	115,4895	184	147.3205	118.4779 149.0331	3.51E-01	6.74E-01	1 Unchanged Low
P28067	D08ab5	major histocompatibility complex;		100	147,3934	125.4983		-3.58E-01	1 Unchanged Low 1 Unchanged Low
Q14126	E11ef7	desmoglein 2	426.8387	366	147,4196		-1.53E+00		1 Unchanged Low
Q14267	P01cd1	regulatory factor X; 4 (influences	127.7912	98.9	147.4351	124.7021		-3.70E-01	1 Unchanged Low
P55197	H09ab2	myelold/tymphoid or mixed-linear	160.668	185	147.4439		-1.24E-01	2.02E-01	1 Unchanged Low
Q9NP87	O20cd8	polymerase (DNA directed); mu	115.1765	86.7	147.4845	116.4497		-4.10E-01	1 Unchanged Low
P04155	A04cd2	trefoil factor 1 (breast cancer; est		136	147.487		-8.07E-01		1 Unchanged Low
Q9C069 P12270	N02gh7	sec13-like protein	52.65015	424	147.5255	207.9658	1.49E+00	3.01E+00	
Q9UIG6	G14cd2 I14ef2	translocated promoter region (to LPAP for lysophosphatidic acid p	104.4493	136 325	147.7447	129,4065	5.00E-01	3.81E-01	1 Unchanged Low
O15453	P08cd5	NBR2	99.08545	108	147.8779 147.9143	230.2685 118.2765	-5.62E-01 5.78E-01	5.72E-01 1.22E-01	1 Unchanged Low
Q9UNW8		G protein-coupled receptor	239.3255	347	148.021	244.7569	-6.93E-01	5.36E-01	1 Unchanged Low 1 Unchanged Low
P06746	110ef6	polymerase (DNA directed); bela	142,5013	225	148.0351	171.8166	5.50E-02	6.58E-01	1 Unchanged Low
O15414	108cd7	trinucleolide repeat containing 4	131.6219	199	148.0947	159.7305	1.70E-01	6.00E-01	1 Unchanged Low
O15259	L22ab7	nephronophthisis 1 (juvenile)	115.0436	110	148.3613	124.437	3.67E-01	-6.59E-02	1 Unchanged Low
Q9NRA8	P05gh4	elF4E-transporter	106.343	178	148.4323	144.3142	4.81E-01	7.45E-01	1 Unchanged Low
Q9NP77 Q06587	C10ef8	HSPC182 protein	171.17	175	148,5234	165.0639	-2.05E-01	3.60E-02	
Q9Y5Z5	P17cd1 E22ef2	ring finger protein 1 heme binding protein 1	119,2377 181,0768	127	148,657	131.6381	3.18E-01	9.12E-02	
015327	105cd4	inositol polyphosphate-4-phosph		196 131	148.7681 148.785	154.5722	-2.84E-01 -3.04E-01	1.13E-01	1 Unchanged Low
P49646	P19cd1	regulator of mitotic spindle assen		117	148.8487	136,7074		-2.95E-01	1 Unchanged Low 1 Unchanged Low
Q99809	F16cd5	conserved gene amplified in oste		91.3	148.9901	124.7728		-5.53E-01	1 Unchanged Low
Q9UBM1	G02cd7	phosphatidylethanolamine N-met	138.4849	122	149.0109	136.3565		-1.88E-01	1 Unchanged Low
P16403	G04ab6	H1 histone family; member 2	427.1258	558	149.1016	377.9287	-1.52E+00	3.84E-01	1 Unchanged Low
075503	B01ab4	ceroid-lipofuscinosis; neuronal 5		75.4	149.1361	124.5021		-9.83E-01	1 Unchanged Low
P15814 Q03164	P20gh6 N01ab7	immunoglobulin tambda-like poly		144	149.2005	123.9169	9.19E-01	8.64E-01	1 Unchanged Low
Q13624	B08ef6	myeloid/lymphold or mixed-lineas interleukin enhancer binding fact	157.0713	81.9 130	149.358	129.4559	-7.26E-02		1 Unchanged Low
Q14118	M02ab5	dystroglycan 1 (dystrophin-assoc		99	149.4265 149.6192	138.6386 118.1383		-6.85E-02 -9.59E-02	1 Unchanged Low
Q13286	E20ab3	ceroid-lipofuscinosis; neuronal 3:	240.521	302	149.6542	230.6187	-6.85E-01	3.27E-01	1 Unchanged Low 1 Unchanged Low
Q9Y5Q0	E15gh7	fatty acid desaturase 3	117.0825	115	149.704	127.2281		-2.72E-02	1 Unchanged Low
Q9P0R1	K08ef2	forkhead box P1	99.17408	114	149.7517	120.8481	5.95E-01	1.96E-01	1 Unchanged Low
P41182	C24ab5	B-cell CLL/lymphoma 6 (zinc fing		85.9	149.853	117.2275		-4.34E-01	1 Unchanged Low
O75497 Q14185	K14cd6	microspherule protein 1	145.7927	163	149.9313	153.0143	4.04E-02	1.64E-01	1 Unchanged Low
095163	M19ab4 P05cd3	dedicator of cyto-kinesis 1 inhibitor of kappa light polypeptic	95.3322	112	150.0398	118.9819	6.54E-01	2.27E-01	1 Unchanged Low
Q14508	L05ef7	WAP four-disulfide core domain:	102.6084	120 201	150.0489 150.0782	157.6357 151.3745	-4.36E-01 5.49E-01	9.73E-01	1 Unchanged Low
P25942	M21ef7	iumor necrosis factor receptor su		239	150.083	166,2403	4.47E-01	1.12E+00	1 Unchanged Low 1 Unchanged Low
Q9Y6E0	F15cd3	serine/threonine kinase 24 (STE:	206,7079	225	150.279		-4.60E-01	1.25E-01	1 Unchanged Low
Q9UQF2	J21cd5	milogen-activated prolein kinase	229.9384	182	150.2902			-3.36E-01	1 Unchanged Low
Q9Y285	D11ab5	phenylalanine-IRNA synthetase-I	179.7613	145	150,3716	158.2291	-2.58E-01	-3.14E-01	1 Unchanged Low
Q04762	F09ab4	cell matrix adhesion regulator	134.1594	176	150.3997	153.4581	1.65E-01	3.90E-01	1 Unchanged Low
Q9Y6J9	P19ef6	TAF6-like RNA polymerase II; p3		158	150,4181	128.6968	9.55E-01	1.03E+00	1 Unchanged Low
O43913 P25208	C06ab8 M05cd1	origin recognition complex; subus nuclear transcription factor Y; ber	105.4862	168	150.4866	155.6272	2.48E-02	1.88E-01	1 Unchanged Low
Q9H9X4	F03gh5	hypothetical protein FLJ11618	189,58	138 179	150.5231 150.5879	131.7807	4.99E-01	3.77E-01	1 Unchanged Low
095297	P17cd4	myelin protein zero-like 1	127.461	121	150.7509	173.1977 133.0095	-3.32E-01 2.42E-01		1 Unchanged Low
Q9UPG8	O06ab8	pleiomorphic adenoma gene-like		165	150.9533	150,4388	1.62E-01	2.95E-01	1 Unchanged Low 1 Unchanged Low
P16333	H16ef5	NCK adaptor protein 1	237.1463	351	150 0510	246.4108		C 00E 04	1 Unchanged Low
Q14108	122ab5	CD36 antigen (collagen type I re-		139		148.3416			1 Unchanged Low
Q9UFF9	005cd5	CCR4-NOT transcription comple:		208	151.0687	174,1041	-1.09E-01	3,55E-01	1 Unchanged Low
015242	O19ab8	nardīlysin (N-arginine dibasic cor		198		186.2788			1 Unchanged Low
Q13442	N15ef3	PDGFA associated prolein 1	156.1376	188		165.0545			1 Unchanged Low
Q99755 Q9NZV1	A15ef6 E09gh2	phosphatidylinositol-4-phosphate cystelne-rich motor neuron 1	112.2177		151.3432		4.32E-01		1 Unchanged Low
Q05195	C09ab7	MAX dimerization protein	136.8996	99.1 254	151.3541	180,6838	4.27E-01		1 Unchanged Low
O9567†	J14ab2	acetylserotonin O-melhyliransfer		98.5	151.5662 151,7064	116.6189	1.47E-01 6.06E-01		1 Unchanged Low
Q9Y658	A09ef2	RNA helicase	112.9715	95,3		120.0494	4.27E-01		1 Unchanged Low 1 Unchanged Low
Q9BTU6	N12gh6	phosphatldylinositol 4-kinase typ		178		142.5367	6.43E-01		1 Unchanged Low
Q9UPN6	G04ef8	KIAA1116 protein	127.681			130,9048			1 Unchanged Low
000170	J03ab2	aryl hydrocarbon receptor interac		223	152.0508	310,1793	-1.87E+00	-1,31E+00	1 Unchanged Low
Q9UJW6	P01ef3	activity-regulated cytoskeleton-as		185		186.4423			1 Unchanged Low
Q9HCT0	N02ef8	fibroblast growth factor 22	157.7471			145,1536			1 Unchanged Low
Q15459 O95400	003cd6 G16cd6	splicing factor 3a; subunit 1; 1201 CD2 antigen (cytoplasmic tail) bir		145		153.5048			1 Unchanged Low
P07954	M06ab3	furnarate hydratase	121.2546	238	152.5257 152.5652		-3.34E-01		1 Unchanged Low
Q9H6Z5	C16gh5	hypothetical protein FLJ21634	101.1467		152.6745		3.31E-01 5.94E-01		1 Unchanged Low 1 Unchanged Low
Q9HCN8	A05gh1	stromal cell-derived factor 2-like				139.1775			1 Unchanged Low
Q92890	A24cd4	ubiquilin fusion degradation 1-lik				176.6159			1 Unchanged Low

043324	N24cd4	eukaryotic translation elongation	150.6748	112	152.9613	138.6164	2.17E-02	-4.25E-01	1 Unchanged Low
Q9NVU8	H10gh2	hypothetical protein FLJ10496	145.4398	119	152.9651	139.1241	7.28E-02	-2.90E-01	1 Unchanged Low
Q9NQW6		anillin; actin binding protein (scra		85.6	152.992			-4.11E-01	1 Unchanged Low
Q9NWH4		milochondrial ribosomal protein t		107			-1.80E-01		1 Unchanged Low
Q9UBV8 P15328	M13cd8 F10ab3	PEF protein with a long N-termin- folate receptor 1 (adult)	152.5051	144 123	153.0828			-4.15E-02	
Q9Y583	B01cd6	NS1-associated protein 1	143.6283	177	153,1163 153,2797	142.7655 157.8847	9.38E-02	-3.14E-01	1 Unchanged Low
Q07326	M08ab8	phosphatidylinositol glycan; class		93	153.3084	126.3257		2.99E-01 -5.12E-01	1 Unchanged Low
075398	M12cd5	SEC22 vesicle trafficking protein-		153		150.1024	9.19E-02		· · · · · · · · · · · · · · · · · · ·
Q9Y2Z8	D09ef2	CGI-09 protein	117.2943	110	153.5282			-9.52E-02	1 Unchanged Low
Q9H3P7	N05gh5	galgi phasphoprotein 1	251.3681	258	153.65	220.9899	-7.10E-01	3.73E-02	1 Unchanged Low
O95678	E03cd5	cytokeratin type II	339.3692	195	153.797		-1.14E+00	-8.00E-01	1 Unchanged Low
P17026	M04cd4	zinc finger protein 22 (KOX 15)	152.726	88.3	153,8779		1.08E-02		1 Unchanged Low
Q9Y3N9 Q9BYT8	M05gh7 · K08gh7	olfactory receptor; family 2; subfa neurolysin (metallopeptidase M3		137	153.9228	139.4653	2.75E-01	1.09E-01	1 Unchanged Low
Q9UHQ7	F02ef8	pp21 homolog	85.99781 119.2248	193 136	154.0005			1.16E+00	
060921	A22ab7	HUS1 checkpoint homolog (S. pc		204	154.126	136.5651 169.0172	3.69E-01 5.29E-02	1.95E-01 4.60E-01	1 Unchanged Low
Q00400	D05ab2	acetyl-Coenzyme A transporter	196.9505	124	154,2012		-3.53E-01	-6.68E-01	1 Unchanged Low 1 Unchanged Low
Q9Y690	H01gh6	mortality factor 4	253.2426	259	154.2307		-7.15E-01	3.23E-02	1 Unchanged Low
Q9NZL9	L24ef7	methionine adenosyltransferase	128.4144	99.7	154.2681	127.4733			1 Unchanged Low
O60563	A05ab4	cyclin T1	179.5706	171	154,2911	168.3854	-2.19E-01	-6.81E-02	1 Unchanged Low
P20591	M03ab7	myxovirus (Influenza virus) resist		873	154.338		-1.57E+00	9,27E-01	1 Unchanged Low
Q9BQC6 P28838	P14gh5 J02ef1	mitochondrial ribosomal protein €		74.3	154.357		-2.89E-01		1 Unchanged Low
Q9NZE5	P08ef1	leucine aminopeptidase 3 hypothetical protein LOC51319	409.1007 135.1502	383 171	154.3828		-1.41E+00		1 Unchanged Low
Q9NX47	K10gh2	hypothetical protein FLJ20445	299.3918	338	154.3904 154.4327	153.3881	1.92E-01 -9.55E-01	3.36E-01	1 Unchanged Low
060612	K11cd2	supervillin	113,628	270	154.4497	179.421	4.43E-01	1.75E-01 1.25E+00	1 Unchanged Low 1 Unchanged Low
O00559	F22cd4	estrogen receptor binding site as		147	154.679		-2.31E-01		1 Unchanged Low
Q9NUU9	E17gh3	hypothetical protein F23149_1	140.3785	224			1.40E-01	6.73E-01	1 Unchanged Low
P22059	C10ab8	exysterol binding protein	133.2924	93.6	154.8714	127.2507		-5.10E-01	1 Unchanged Low
000408	105ef6	phosphodieslerase 2A; cGMP-stl		169	154.9688		-2.47E+00		1 Unchanged Low
O94935 P41134	P22ef7	SAC1 suppressor of actin mutatic		128	154.9914	142,9113			1 Unchanged Low
P24407	H15ef5 M08ab7	inhibitor of DNA binding 1; domir mel transforming oncogene (dert		292 189	155.0439		-1.49E+00		1 Unchanged Low
O00232	N05ab8	proteasome (prosome; macropali		139	155.1684 155.2465	189.5598 139.0276	-5.30E-01 3.42E-01		1 Unchanged Low
Q01432	G04ab2	adenosine monophosphate dean		117	155,341	127.7664	4.83E-01	1.85E-01 7.10E-02	1 Unchanged Low 1 Unchanged Low
Q13525	G15cd4	Interleukin 1 receptor-like 2	136.0802	130	155.4674	140.5051		-6.63E-02	1 Unchanged Low
Q16611	N04ab3	BCL2-antagonist/killer 1	280.4131	194	155.5414		-8.50E-01		1 Unchanged Low
Q9NPA3	C11gh5	hypothetical protein STRAIT1149		85.5	155.5531	117,8671		-3.96E-01	1 Unchanged Low
Q9GZM8	D14gh7	LIS1-Interacting protein NUDEL;		131	155.5611	141.4354		-7.41E-02	1 Unchanged Low
P42702 Q05940	P06ab6 P07ef5	leukemia inhibitory factor recepto		112	155.6189	120.0363	7.51E-01	2.77E-01	1 Unchanged Low
Q9Y3Q5	H18cd6	solute carrier family 18 (vesicular putative DNA/chromatin binding)		153 150	155.637 155.6726	127.0621	-1.27E+00 1.04E+00		1 Unchanged Low
Q15756	H08ab7	potassium inwardly-rectifying cha		60.9	155.7504	128.583		9.77E-01 -8.81E-01	1 Unchanged Low 1 Unchanged Low
075386	M17cd3	tubby like protein 3	128.6914	82.7		122.3782		-6.39E-01	1 Unchanged Low
P50750	D03ef5	cydln-dependent kinase 9 (CDC:	270.8229	215			-7.97E-01		1 Unchanged Low
Q9NWY6	A18gh3	hypothetical protein FLJ20531	91.85185	130		125.9903	7.65E-01	5.01E-01	1 Unchanged Low
P50391	K05cd1	pancreatic polypeptide receptor	136.5278	112	156.1953	134.9143		-2.85E-01	1 Unchanged Low
Q9NX07 O75354	L20gh6 A15ab4	tRNA selenocysteine associated ectonucleoside triphosphate dipt	258,434 386,4989	287 258	156.2449		-7.26E-01	1.52E-01	1 Unchanged Low
Q9H832	L12gh5	hypothetical protein FLJ13855	173.4437	307	156.4638 156.9292		-1,30E+00		1 Unchanged Low
Q9UBW8		COP9 constitutive photomorphog		111	157.0119	123.4686	-1.44E-01 6.11E-01	8.22E-01 1.05E-01	1 Unchanged Low 1 Unchanged Low
P14136	E23ab5	glial fibrillary acidic protein	108,166	86.6	157,2761	117.3353		-3.21E-01	1 Unchanged Low 1 Unchanged Low
Q9NYF3	F08ef1	chromosome 5 open reading fran	121.4116	94.3		124,4392		-3.64E-01	1 Unchanged Low
Q15056	A09gh2	Williams-Beuren syndrome chror	202.4969	124	157.6783	161.3602	-3.61E-01		1 Unchanged Low
O15534	116ab8	period homolog 1 (Drosophila)	237.2007	143	157.7195	179,3665	-5.89E-01		1 Unchanged Low
Q9Y399 O14972	O21ef2	mitochondrial ribosomal protein \$		139.	157.7339		-5.44E-01		1 Unchanged Low
Q14113	AD4cd6 M15ab2	Down syndrome critical region ge AE binding protein 1	124.9796	267	157.7386	218,3678	-5.45E-01	2.16E-01	1 Unchanged Low
O95755	O18cd5	RAB36; member RAS oncogene	132.9919	95.3 97	157.955 158.0474	126.0667 129.3518		-3.92E-01 -4.55E-01	1 Unchanged Low
Q9NPC3	109gh4	enhancer of invasion 10	104.2128	121	158,1023	127.8011	6.01E-01	2.17E-01	1 Unchanged Low 1 Unchanged Low
Q9Y6E9	N24cd7	siduln silent mating type Informal		153		140.9006	5.00E-01	4.48E-01	1 Unchanged Low
Q9Y5B8	D17cd8	NME7	92.69165	116	158.2444		7.72E-01		1 Unchanged Low
P49642	121ab8	primase; polypeplide 1 (49kD)	135.2376	70.2	158.2887	121.2327	2.27E-01	-9.47E-01	1 Unchanged Low
O60660 Q9H251	J04ab2	ash2 (absent; small; or homeotic	274.3502	177			-7.93E-01		1 Unchanged Low
Q9Y6C2	008gh7 A22cd7	cadherin related 23 elastin microfibril Interface locate	113.1038	184	158.4524	151.9031		7.03E-01	1 Unchanged Low
O95825	D13cd5	crystallin; zeta (quinone reductas		92.2 133	158.6648 158.6719	119.6097		-2.28E-01	1 Unchanged Low
Q9UHG3	J02ef2	prenylcysteine lyase	250.3969	207	158.8419	145.1487	-6.57E-01	-1.17E-01	1 Unchanged Low 1 Unchanged Low
000629	J18ab6	karyopherin alpha 4 (Importin alp		295	158.8458		-1.71E+00		1 Unchanged Low 1 Unchanged Low
Q15397	B14gh1	KIAA0020 gene product	109.8533	134		134.3527		2.89E-01	1 Unchanged Low
Q14999	H01gh1	KIAA0076 gene product	71.35119	173	159.0037	134.6153	1.16E+00		1 Unchanged Low
Q12772	M18cd2	sterol regulatory element binding		94.6	159.1986	137,4667	5.34E-03	-7.46E-01	1 Unchanged Low
075150	F09gh1	ring finger protein 40	129.2351	169		152,3442		3.84E-01	1 Unchanged Low
043423 043311	L07gh6 D12cd8	acidic (leucine-rich) nuclear phos		143	159.4372	147.5392		2.63E-02	1 Unchanged Low
Q9NYX4	D12cg8 B07ef4	muscleblind-like (Drosophila) calcyon; D1 dopamine receptor-ii	165.2243	99.4	159,4681	141,3508	-5.12E-02		1 Unchanged Low
Q9HD71	G04gh4	hypothetical nuclear factor SBBI:		147 140	159.5302 159.5491		-3.78E-01	1.82E-01	1 Unchanged Low
Q9NZH2	N03cd8	replication initiation region protei		179			-4.96E-01		1 Unchanged Low 1 Unchanged Low
Q9NVH1	B05gh3	hypothetical protein FLJ10737	137.4673	217	159.6304	171.2625			1 Unchanged Low
Q9NVG3	B11gh3	hypothetical protein FLJ10751	96.28296	105	159.6829		7.30E-01	1.19E-01	1 Unchanged Low
P52179	A21cd4	myomesin 1 (skelemin) (185kD)	157.0608	94	159.8272	136.9676	2.52E-02	-7.40E-01	1 Unchanged Low
094815	B09cd3	frizzled homolog 1 (Drosophila)	93.97877	101	159.9198	118.2939	7.67E-01	1.04E-01	1 Unchanged Low
Q9UBY9 Q9NVX7	G01ef4 G18gh3	heat shock 27kD protein family; r hypothetical protein FLJ10450		102	160.1725	120.3216	6.94E-01	4.08E-02	1 Unchanged Low
095704	A02cd6	FE65-like protein 2	111.5196 99.35088	121	160.2507	130.905	5.23E-01	1.17E-01	1 Unchanged Low
	+ • •	me present e	20.03000	113	160.3239	141.1913	3.500-01	2.06E-01	1 Unchanged Low

D40004										
P19801	A23ab2		in 164.6296	68.	3 160,3477	7 131.0836	-3.80E-02	-1.27F+00		Unchanged
043709		Williams Beuren syndrome chr	or 127.8538	13		140.7865				Unchanged Low
Q12839		H326	109 677	110				8.26E-02		Unchanged Low
043503		RAD51 homolog C (S. cerevisi	€ 96.34779	97.						Unchanged Low
Q07898		CO163 antigen	87.73563	177						Unchanged Low
O96015		dynein; axonemal; light polyper	11 94 54055	25				1.01E+00		Unchanged Low
Q92524	L11ab8	proleasome (prosome; macrop	ali 396 6515	220			7.08E-01	1.41E+00		Unchanged Low
Q92692	B06ab8	pollovirus receptor-related 2 (h	or 302 7105				-1.30E+00		1	Unchanged Low
Q9Y6Y8		Sec23-Interacting protein p125		317				6.60E-02	1	Unchanged Low
Q9NZ78		uncharacterized bone marrow p	104.2429	131			6.30E-01	3.34E-01	1	Unchanged Low
Q15120		manufo dobudences litera	n 107.2001	127			5.90E-01	2.41E-01		Unchanged Low
P35241	F22ab8	pyruvate dehydrogenase kinasi		107				1.43E-01	1	Unchanged Low
Q9Y326	D23ef2	nakin	276.6012	213	3 161.7003	216.9693	-7.74E-01	-3.80E-01	1	Unchanged Low
Q92600		brain specific protein	359,3582	160	161.7123		-1.15E+00		•	Unchanged Low
000487	J03cd5	RCD1 required for cell different	ia 115.6633	145	161.7142					Unchanged Low
	N14cd5	26S proteasome-associated pa		275	181.8834		-1.02E+00		i	
Q9Y689	P21cd7	ADP-ribosylation factor-like 5	137.7737	159	161.8965	152,9706	2.33E-01	2.09E-01		Unchanged Low
Q9ULB8		CMP-NeuAC:(beta)-N-acetylgal	a 91.01128	121			8.31E-01	4.09E-01		Unchanged Low
O60897	L09cd7	RAB; member of RAS oncogena	89.32474	112			8.59E-01		1	
P51809	J05cd2	synaptobrevin-like 1	228.3546	141			-4.94E-01	3.226-01		Unchanged Low
075581	A03ab7	low density lipoprotein receptor	122,4708	125					1	
043795	F12ef4	myosin class I; myh-1c	259,4543	282			4.05E-01	2.53E-02	1	Unchanged Low
Q14188	H17ef5	transcription factor Dp-2 (E2F d	ir 107.308					1.20E-01	1	Unchanged Low
P06280	B07ab3	galactosidase; alpha	178.0238	204			5.96E-01	9.25E-01	1	Unchanged Low
Q14541	O17ab7	hepatocyte nuclear factor 4; gar	1/0.0236 - 540.0007	304				7.70E-01	1	Unchanged Low
P57723	E17gh8	poly(rC) binding protein 4		283		331.895	-1.76E+00	-9.57E-01	1	Unchanged Low
	3 A24gh3	DEADAL (Acc Clu Ale Accents	188,2845	231			-1.97E-01	3.08E-01		Unchanged Low
Q9NPC5		DEAD/H (Asp-Glu-Ala-Asp/His)	154,71	104			7.22E-02	-5.71E-01		Unchanged Low
Q9Y3C5	B19e/3	nudix (nucleoside diphosphate)		98.4	162.6929	122.7442	6.02E-01	-1-24E-01		Unchanged Low
	B01gh2	ring finger protein 11	160.0631	164	162.7518	162,3366	2.40E-02			Unchanged Low
		chromosome 20 open reading tr	z 98.58482	95.6	162.7744	118.9844	7.23E-01			Unchanged Low
P52742	122cd3	zinc finger protein 135 (clone pl	ł 81.4141	121				5.68E-01		
Q9Y313	D17ef2	PTD013 protein	90.25232	116	163,4053	123,3545	8.56E-01	3.67E-01	:	Unchanged Low
Q9P2W9		syntaxin 18	128.6426	188	163.4277	160.0555	3.45E-01	5.48E-01		Unchanged Low
P14778	F09ab6	interleukin 1 receptor; type I	264.5401	539	163,4432				1	Unchanged Low
Q15464	l12ef5	SHB (Src homology 2 domain or	284.291	301				1.03E+00		Unchanged Low
015444	H07cd2	small inducible cytokine subfami	244,6426	97.1	163,6053			8.24E-02		Unchanged Low
P49796	P14ef4	regulator of G-protein signalling	220.2769	99.3		168.4558	-5.80E-01 ·	1.33E+00		Unchanged Low
Q9NUS7	B04gh3	hypothetical protein FLJ11164	81.76063	114	163.6422	161.0774				Unchanged Low
P48651	Lttef3	phosphalidylserine synthase 1	211.8695	139				4.77E-01	1	Unchanged Low
Q07706	F15cd4	tetracycline transporter-like prote	1/3 0701		163,6792 163,7206		-3.72E-01		1	Unchanged Low
075143	B11gh1	KIAA0652 gene product	106.5331	144		150.2019		7.31E-03	1	Unchanged Low
075039	L19gh1	KIAA0451 gene product		155	163.7687	141.6405		5.37E-01	1	Unchanged Low
P02261	M10gh6	H2A histone family; member N	174.142	135	163.8572		-8.78E-02		1	Unchanged Low
Q9H9A0	L20gh5	hypothetical protein FLJ12895	93.48591	273	163.8621	176.9259	8.10E-01	1.55E+00		Unchanged Low
Q9NV35	J21gh3	hypothetical protein FLJ10956	86.42599	130	184,3655	126,7707		5.84E-01		Unchanged Low
Q9P1G8	C05gh4	WW domain containing the start	99.15342	154	164.4852	139.1489	7.30E-01	6.33E-01		Unchanged Low
P26368	D11cd7	WW domain-containing adapter	119.4436	182	164.5311	155.3256	4.62E-01	6.08E-01		Unchanged Low
Q98SZ7	E11gh8	U2 small nuclear ribonucleoprote	217.0564	107	164.8861	163.126	-3.97E-01 -	1.01E+00	11	Unchanged Low
Q9Y391	G23ef2	DEAD/H (Asp-Glu-Ala-Asp/His) i	130.7858	111	164.9084		3.34E-01 -		1	Unchanged Low
Q9NYR5	E05ef3	androgen-regulated short-chain	222.3497	128	164.9823		4.31E-01			Unchanged Low
P05165		HSPC126 protein	99.37243	110	165.2098			1.46E-01		Inchanged Low
Q9NPF4	N12ab7	propionyl Coenzyme A carboxyla	100,892	120	165,6902		:	2.54E-01	11	Unchanged Low
	021gh3	O-sialoglycoprotein endopeptida	103.8223	112	165.9716			1.13E-01		Unchanged Low
P16106	D01cd3	H3 histone family; member K	99.19006	130			- :	3.87E-01	1 1	Unchanged Low
P3\$249	003ef5	replication factor C (activator 1)	105.2406	98.7			6.59E-01 -		- 13	Inchanged Low
P28702	P18gh1	retinoid X receptor; beta	74.914	142		127.7227				Unchanged Low
P49674	K14ab4	caseln kinase 1; epsilon	99.33956	145					1 1	Inchanged Low
Q9NXW2		DnaJ (Hsp40) homolog; subfamil	247,7907	235		216.5041 -	F 745 04	5.44E-01		Inchanged Low
014672	101ab2	a disintegrin and metalloproteina	378.0042	226	168.5417	210.3041	5.74E-U1 ·	7.47E-02	1 (Inchanged Low
Q9Y5B9	120cd7	chromatin-specific transcription e	163 0543	188		256.8185 -1			1 8	Inchanged Low
Q9H9J2	H22gh5	mitochondrial ribosomal protein I	206,428	256				2.04E-01	1 L	Inchanged Low
P23759	G06ab8	paired box gene 7						3.10E-01	1 (Inchanged Low
Q16643	L05ab4	drebrin 1	92.41833 265.008			153.0178	8.52E-01 1	-11E+00	1 (Inchanged Low
P48454	121cdt	protein phosphatase 3 (formerly:	10E 040C			194.7352 -			1 (Inchanged Low
Q9Y508	F21ef4	zinc finger protein 313				147.2674	6.57E-01	6.72E-01	1 (Inchanged Low
	G09cd2	Singren syndrome antique A4 /5'	331,595	395	167.2355	297.9139 -	9.88E-01 (2.52E-01		Inchanged Low
	N13ab5	Sjogren syndrome antigen A1 (5) v-crk sarcoma virus CT10 oncogi	301.1776	183	167.414	217.1349 4	B.47E-01 -	7.20E-01	1 1	Inchanged Low
	B19cd4	UDP abroadtractomes 2 feet	131.2065		187.7716	153.202	1.50E-01 -	1.05E-01	1 1	Inchanged Low
044040	H14cd1	UDP glycosyltransferase 2 family				144.2152	2.83E-01 -1	1.20E-01		Inchanged Low
		selectin P ligand	112.9679	130	167.9184	136,8261	5.72E-01	1.98E-01		
	P17gh8	hypothetical protein FLJ14525	86.75908	113	167.974		9.53E-01			Inchanged Low
	L23ef1	homolog of yeast exosomal core		140			1.44E-01 -1		4 1	Inchanged Low
	N03cd1	somatostatin receptor 4	147.3942				1.89E-01 -6		1 0	Inchanged Low
	003ab4	diptheria toxin resistance protein	210.9658			183.0009 -	7.00E-01 *C	125.04		nchanged Low
	L24ab8	plakophilin 2	362 0423			247.4823 -1	410,00	1 12E-U1		nchanged Low
	B21gh7	leukocyte immunoglobulin-like re	139 2571	234	168.1829	180 4876	7700 04	.00E-01	1 U	nchanged Low
	P18ab5	down-regulator of transcription 1	205 2249		168,2606	180.4876 2	ZE-01 7	.49E-01		nchanged Low
Q9HCS4	P10gh7	HMG-box transcription factor TCI	76.48422		100,2000	165.0596 -2	4.67E-01 -7	.54E-01	1 U	nchanged Low
Q9P035	N20ef2	butyrate-induced transcript 1	102.716		168.2926	133.5936 1	.14E+00 1	.03E+00	1 U	nchanged Low
	P24gh3	amyotrophic lateral sclerosis 2 (in			168.3034	137.5999 7	7.12E-01 4	.65E-01		nchanged Low
	13ef3	chromosome 5 open reading fran	102,352			126.1345 7	'.19E-01 7	.24E-02		nchanged Low
		chromosome 21 open reading fra	09.7.2300		168.4424	131.7509 g	0.09E-01 6	.11E-01		nchanged Low
	K16cd6	glioma amplified on chromosome	448 2000		168.5171	121.6076 7	'.70E-01 -2	.03E-02		nchanged Low
		transient receptor palacitat	110.3239		168.5172	122.705 5	.35E-01 -4	.82E-01		nchanged Low
	14ef8	transient receptor potential cation				118.5764 B	1.06E-01 -8		1 11	nchanged Low
		KIAA1084 protein	101.3235	111	168.5336	126,797 7	345.01 1	26E-01	1 11	nchanged Low
	21er4 024cd1	G protein-coupled receptor kinas		167	168.7352 1	165.5706 7	.00E-02 5	.72E-02		nchanged Low
	724CU I	secreun receptor	174 EE72	209	168./38 1	164.1802 🎿	98F-02 2	60F-01		
	211W9	receptor-Interacting serine-threor		388	168,//22 2	2/6,5664 .e	Q1E_01 5	11F-01		nchanged Low
P32321 F	214ab\$	dCMP deaminase	218.5685	60.1	168.8404 1	149.1554 -3	725.01 4	REE+DO		nchanged Low
									1 0	nchanged Low

	J16ab4	hypocretin (orexin) neuropeptide			168.9064 168.9596	143.2577 177.8201			1 Unchanged Low 1 Unchanged Low
	M15ab4 115ef8	deoxyribonuclease Hike 2 PRO0038 protein	218.1787 89.17218		168.9849		9.22E-01		1 Unchanged Low
P30519	N05ab6	heme oxygenase (decycling) 2	586.3897		169.1269	380,9032 -	1.79E+00	-5.99E-01	1 Unchanged Low
Q9NVE7	D03gh3	hypothetical protein FLJ10782	141.6695		169.2644	152,9293	2.57E-01	6.16E-02	1 Unchanged Low 1 Unchanged Low
	A14ef8	HSPC154 protein bromodomain-containing 7	227.9143 147.6635		169.4393 169.5045	214.317 139.9456	-4.28E-01 1.99E-01	1.08E-01 -5.24E-01	1 Unchanged Low
	M04cd8 E01cd6	spermatogenesis associated 2	230.7114		169.6678		-4.43E-01	-6.20E-02	1 Unchanged Low
	N14ef1	zinc finger protein 219	105.6753		169.7666		6.84E-01	7.27E-01	1 Unchanged Low
	H19gh7	recombination protein REC14	124.7245		169.9984	152.3724 174.0094	4.47E-01 -3.52E-01	3.81E-01 -6.85E-01	1 Unchanged Low 1 Unchanged Low
P07686 Q9NZV6	J17ab3 L06ef2	hexosaminidase B (bela polypep selenoprotein X; 1	241.3633		170.0092 170.0232		-5.05E-01	-2.60E-01	1 Unchanged Low
014678	E17cd1	ATP-binding cassette; sub-family			170.0525	124.9148	1.13E+00	7.10E-01	1 Unchanged Low
P29972	O04ab2	aquaporin 1 (channel-forming int-	309.1218		170.2078			-1.21E+00	1 Unchanged Low 1 Unchanged Low
095373	D03cd6	RAN binding protein 7 Sec23 homolog A (S. cerevisiae)	247.4368	113 100	170.45 170.9386			-1.13E+00 1.52E-01	1 Unchanged Low
Q15436 Q9BQ67	O12cd6 K23gh8	glutamate rich WD repeat proteir		112	170.966	141.0924		-3.25E-01	1 Unchanged Low
P35321	D21gh6	small proline-rich protein 1A	62.28271		171.0104			1.93E+00	1 Unchanged Low
Q9UPQ8	110ef8	KIAA1094 protein	76.76245 95.36294	123 92.4	171.0467 171.0704	123.4833 119.6038	1.16E+00	6.76E-01 -4.59E-02	1 Unchanged Low 1 Unchanged Low
O43292 Q9UBM3	A19cd4 G02cd1	GPAA1P anchor attachment prot proteasome (prosome; macropali			171.1554		-7.49E-01	-4.93E-01	1 Unchanged Low
P19338	M20ab7	nucleolin	148.8642		171.2094	135.8947		-7.65E-01	1 Unchanged Low
Q9NPE6	E17gh1	sperm associated antigen 4	111.172		171.4356	139.3341	6.25E-01		1 Unchanged Low
000442	J12cd3	RTC domain containing 1	274.6701	279 108	171.531 171,5949	241.6967 120.5873	-6,79E-01		1 Unchanged Low 1 Unchanged Low
P28356 P49447	D09ef1 G06ef1	homeo box D9 cytochrome b-561	84,5638 174,7152	167	171.6447			-6.39E-02	1 Unchanged Low
Q9UQA4	P18cd4	nuclear factor (erythroid-derived	227,4253	470	171.8945	269.7702			1 Unchanged Low
015254	D09ab2	acyl-Coenzyme A oxidase 3; pris		192	172.0218				1 Unchanged Low
Q9NWW8		hypothetical protein FLJ20559	198.7861 154.1159	220 140	172.0261 172.0658	155.2376	-2.09E-01	1.48E-01 -1.43E-01	1 Unchanged Low 1 Unchanged Low
O75391 Q92959	M08cd5 H19cd2	sperm associated antigen 7 solute carrier family 21 (prostagli		102	172.112		9.97E-01		1 Unchanged Low
000499	G06ab2	bridging integrator 1	162.8584	111	172.4453	148.8447		-5.50E-01	1 Unchanged Low
Q14197	J05ab6	immature colon carcinoma transc		102	172.5526	121.725	9.24E-01		1 Unchanged Low 1 Unchanged Low
Q9NXS9	K15gh2 N18cd6	hypothetical protein FLJ20071 tetraspan transmembrane 4 supe	121,4575 83,06795	103 119	172.5552 172.5557	132,3749 124,8565	1.05E+00	-2.36E-01 5.18E-01	1 Unchanged Low
O75954 Q9UJY9	E17cd8	lifeguard	156,5909	75.6	172.6495	134.9529		-1.05E+00	1 Unchanged Low
P82912	F10gh5	mitochondrial ribosomal protein 5		196	172.7091	158,0886	7.17E-01		1 Unchanged Low
Q12918	J02ab6	killer cell lectin-like receptor subl		290 194	172.8141 173.0059			4.97E-01 1.01E+00	1 Unchanged Low 1 Unchanged Low
Q14789 Q04206	E01ab6 123ef6	golgi autoantigen; golgin subfam v-rei reticuloendotheliosis viral o		242	173.0443			-4.40E-01	1 Unchanged Low
P19838	M15ef5	nuclear factor of kappa light poly		277	173.0997		-4.05E-01		1 Unchanged Low
Q92748	A24cd2	thyroid hormone responsive (SPC		88.2	173.314			-2.02E-01	1 Unchanged Low 1 Unchanged Low
O95095	P13ab2 K18ab8	amyloid beta precursor protein (c phosphogluconate dehydrogenate		201 147	173.3314 173.42	191.35 177.5327	-2.06E-01		1 Unchanged Low
P52209 P42566	H10ef5	epidermal growth factor receptor		158	173.4643		-1.68E-01		1 Unchanged Low
Q13772	D05gh6	nuclear receptor coactivator 4	273.5167	367	173,5038		-6.57E-01		1 Unchanged Low
P43897	D09gh6	Ts translation elongation factor;		226 86.2	173,8489 174,5738			-5.28E-01 -2.69E-01	1 Unchanged Low 1 Unchanged Low
P09329 O7561B	D08ef7 F24cd4	phosphoribosyl pyrophosphate s death effector domain-containing		176	174,7689				1 Unchanged Low
Q9UHI9	B18ef1	flavohemoprotein b5+b5R	169.2028	197	174,7919				1 Unchanged Low
Q9H824	J07gh6	F-box only protein 22	89.2751	161	175.62			8.50E-01 2 -1.64E-01	1 Unchanged Low 1 Unchanged Low
Q9UQ43	G07ab3 N08ef1	BAI1-associated protein 2 sialyltransferase 8C (alpha2;3Ga	184.8198 220.9909	165 256	175,6399 175,9869				1 Unchanged Low
O43173 Q9BZZ2	G13gh7	sialoadhesin	147.52	189	175.9952				1 Unchanged Low
Q92611	116gh1	KIAA0212 gene product	96,3144	94.3	176.0348			-3.08E-02	1 Unchanged Low
Q969V6	K14gh7	megakaryoblastic leukemia (tran		161 106	176.0531 176.1145			1 2.49E-01 1 -5.38E-01	1 Unchanged Low 1 Unchanged Low
P08107 Q9BQ24	K14ab7 E21gh6	heat shock 70kD protein 1B hypothetical protein MGC2550	153.5141 153.9741	255	176,1143				1 Unchanged Low
Q9H7X1	N07gh5	hypothetical protein FLJ14153	155.291	236	176.5938				1 Unchanged Low
Q99616	D09cd2	small inducible cytokine subfami		100	176.5958			1 -3.79E-01	1 Unchanged Low 1 Unchanged Low
Q92896	L04cd7	golgi apparatus protein 1 Mad4 homolog	97.73429 210.3185	90.3 244	176,622 176,6383		8.54E-0 2.52E-0 -		1 Unchanged Low 1 Unchanged Low
Q14582 Q13505	J16ef6 K21ab7	metaxin 1	199.1231	168	176.6745			1 -2.46E-01	1 Unchanged Low
Q13722	O24cd8	chromosome 3p21.1 gene seque		144	176.7101	135.6336	1.04E+0	7.49E-01	1 Unchanged Low
P01308	110ab6	Insulin	367.022	113	176.7437	7 218.8465 3 238.6383	-1.05E+0	0 -1.70E+00 1 3.92E-01	1 Unchanged Low 1 Unchanged Low
O00405 P41215	D19ab8 E03ab5	prolein phosphatase 1; regulator fatty-acid-Coenzyme A ligase; lo		306 199				1 -4.27E-01	1 Unchanged Low
P31314	F04ef6	homeo box 11 (T-cell lymphoma		170					1 Unchanged Low
095427	G21cd8	phosphatidylinositol glycan; clas	84,22817	116			1.08E+0		1 Unchanged Low
Q12981	C24ef6	BCL2/adenovirus E1B 19kD Inte		99.8 184	177.5246		1.11E+0	0 2.77E-01 1 -2.03E-01	1 Unchanged Low 1 Unchanged Low
Q9Y5V1 Q9H4B4	L10ef1 022ab4	HSPC042 protein cytokine-inducible kinase	212.074 161.982	194					1 Unchanged Low
Q9H6D1				108	177.8526	126.9312	8.98E-0	1 1.72E-01	1 Unchanged Low
O60504	J18cd5	vinexin beta (SH3-containing ad		99.8				1 -1.33E-01	
Q9UNK0		syntaxin 8 proline-serine-threonine phosph	218,2149	171 158				1 -3.53E-01 0 -1.23E+00	
O95657 P20645	B24cd4 C05ab7			234				1 -6.03E-01	1 Unchanged Low
014569	B03ab2	putative tumor suppressor	122,4884	79.2	178.13	126.62	2 5.40E-0	1 -6.28E-01	1 Unchanged Low
015293	D18cd3	MAP-kinase activating death do		114				1 -4.59E-01	
P00390	E14ef7 K24gh5	glutathione reductase PP3111 protein	282.1492 229.2714	196 361		3 256.1517		1 -5.24E-01 1 6.54E-01	
Q9H8V3 Q14558				227				1 -4.09E-01	1 Unchanged Low
014710	F01ef5	cell cycle progression 2 protein	261,2178	189	178.792	3 209.710	3 -5.47E-0	1 -4.66E-01	1 Unchanged Low
Q9UJ70	J10ef3	N-acetylglucosamine kinase	151.2986	276		2 201.938 [.] 8 226.589;	1 2.41E-0		
Q13397 Q9NRG	M18cd6 4 B21gh4		215.883 143.1326	265 279		9 200,333			
GRIANG	, DEIGHT	w preside		_,,					-

251312	M	DID III I							0.20
P51449	B01cd2	RAR-related orphan receptor C	236.6977	97.2	179.0372		-4.03E-01		1 Unchanged Low
Q9H768	O21gh7	synaplotagmin-like 2	83.78255	121	179.0813	128.0355	1.10E+00	5.33E-01	1 Unchanged Low
Q14332	F22ef5	frizzled homolog 2 (Drosophila)	86.25011	129	179.1323	131,3078	1.05E+00	5.76E-01	1 Unchanged Low
Q13207	N13cd2	T-box 2	118.9378	108	179.1858	135,2524	5.91E-01	-1.44E-01	1 Unchanged Low
Q9H6B8	B19gh5	hypothetical protein similar to mo	134.338	129	179.5015	147.5119	4.18E-01	-6.19F-02	1 Unchanged Low
000462	J09ab7	mannosidase; bela A; lysosomal	83,23817	107	179,6953	123.2365	1.11E+00		1 Unchanged Low
Q9Y3DB	H14ef2	adrenal gland protein AD-004	178.4931	163	179.7989	173.7578	1.05E-02		1 Unchanged Low
P17900	F19ab3			232					
Q9P0M4	O10cd8	GM2 ganglioside activator proteir			179,914	178,9056	5.28E-01		1 Unchanged Low
COLOMA		inlerleukin 17C	260.8753	223	180.0241		-5.35E-01		1 Unchanged Low
0011700	F13gh8	G protein-coupled receptor 54	87.92591	133	180.0735	133.5723		5.94E-01	1 Unchanged Low
Q9NZ92	J02gh3	SWI/SNF related; matrix associa		247	180.208	231.4963	-5.71E-01		t Unchanged Low
Q9NX94	L18gh6	hypothetical protein FLJ20154	133.6318	165	180.3424	159.7713	4.32E-01	3.07E-01	1 Unchanged Low
P04049	G12ef5	v-raf-1 murine leukemia viral onc	225.086	183	180.3925	196.2063	-3.19E-01	-2.98E-01	1 Unchanged Low
Q13181	B06cd1	ATP-binding cassette; sub-family	167.9383	127	180,4651	158,5133	1.04E-01	-4.02E-01	1 Unchanged Low
P22102	M05ab6	phosphoribosylglycinamide form	295.0489	180	180.5026	218,4202	-7.09E-01	-7.15E-01	1 Unchanged Low
Q9NQZ8	A09gh5	endothelial zinc finger protein Inc		138		138,4428	9.01E-01		1 Unchanged Low
Q9UN30	D04cd2	sex comb on midleg-like 1 (Drost		86	181.0121	121.4921	8.93E-01		1 Unchanged Low
P26651	J04ef6	zinc finger protein 36; C3H type;		162	181.0638	209,8211	-6.60E-01		
P18887	006ef6	X-ray repair complementing defe		102				2.54E-01	1 Unchanged Low
Q9Y2S0	P14ef1				181.0778	122.9254			1 Unchanged Low
		RNA polymerase I 16 kDa subun		149	181,2184		-1.36E-02		1 Unchanged Low
P49023	105ef7	paxillin	313.7788	334	181.2995		-7.91E-01	9.00E-02	1 Unchanged Low
P49321	O03ab7	nuclear autoantigenic sperm prof		78.2	181.3524	126.4866	5.97E-01		1 Unchanged Low
Q13428	M14cd1	Treacher Collins-Franceschelti s		123	181.4282	156,2838	1.43E-01	-4.18E-01	1 Unchanged Low
Q9HCS6	G07gh4	vacuolar protein sorting 11 (yeas	177.2833	302	181.4601	220.0848	3.36E-02	7.66E-01	1 Unchanged Low
P20813	102gh6	cytochrome P450; subfamily IIB (122.3562	162	181.476	155.3947	5.69E-01	4.08E-01	1 Unchanged Low
000459	C05cd1	phospholnositide-3-kinase; regul	145.3741	160	181.534	162,3275	3,20E-01	1.39E-01	1 Unchanged Low
P50151	H24ab4	guanine nucleotide binding prote	186,5329	193	182.0963	187.0796	-3.47E-02	4,62E-02	1 Unchanged Low
Q16850	D12ab3	cytochrome P450; 51 (lanosterol		116	182.1309		-9.59E-01		1 Unchanged Low
Q9ULW3		TATA-binding protein-binding pro		207	182,1565	162.0616	9.06E-01		1 Unchanged Low
Q16378	O20cd7	proline rich 4 (lacrimal)	81.91	152	182.3589	138,8678		8,95E-01	1 Unchanged Low
P41240	J14ab4	c-src lyrosine kinase	175.1285	153	182.4257	170.0806			1 Unchanged Low
Q16649	G05cd1						5.89E-02		
Q9Y3M2		nuclear factor; interleukin 3 regul		169	182.4266		-9.55E-01		1 Unchanged Low
	J03ef8	chromosome 22 open reading fra		194	182.6022	150.2264			1 Unchanged Low
Q9UK58	M14gh4	cyclin L ania-6a	191,3619	313	182.6351		-6.73E-02		1 Unchanged Low
P47874	M23cd1	olfactory marker protein	250,3691	172	182.73	201.6274	-4.54E-01		1 Unchanged Low
095620	O03cd7	protein similar to E.coli yhdg and	227,8932	207	183,1055	205,8925	-3.16E-01	-1.41E-01	1 Unchanged Low
CAHN5	J11cd8	CDK4-binding protein p34SEI1	144.2806	163	183.5211	163,6559	3.47E-01	1.77E-01	1 Unchanged Low
P02810	F07gh6	proline-rich protein Haelll subfan	410,3478	392	183.6977	328,6668	-1.16E+00	-6.62E-02	1 Unchanged Low
094903	K04cd7	proline synthetase co-transcriber	242,2062	146	183.9744	190,7733	-3.97E-01	-7.29E-01	1 Unchanged Low
Q9P0W6	103ef4	lipopolysaccharide specific respo		177	184.0725	169.9232	3,04E-01		1 Unchanged Low
Q9H8X4	O15gh5	stromal membrane-associated pr		342	184.0879		-9.60E-01		1 Unchanged Low
P02786	D24ef5	transferrin receptor (p90; CD71)	320,8365	257	184.2545		-8.00E-01		1 Unchanged Low
	E15ef1	immunoglobulin heavy constant (176	184.3533		-1.07E+00		1 Unchanged Low
Q99717	J01ab7	MAD; mothers against decapents		153	184.3978		-3.04E-01		1 Unchanged Low
Q15036	B23gh1	sorting nexin 17	303,8909	411	184,4831			4.34E-01	
P31751	A08ab2	v-akt murine lhymoma viral onco		253	184.8245		-4.57E-01		1 Unchanged Low
P11177	G21ab8	pyruvate dehydrogenase (lipoam		150	185.0719				1 Unchanged Low
Q9UEQ6	F22ab2	arrestin; beta 2	190,7331	128			-3.08E-01		1 Unchanged Low
Q16531	J11ab5				185.1221		-4.31E-02		1 Unchanged Low
P28698		damage-specific DNA binding pn		229	185.1363		-1.48E-01	1.60E-01	1 Unchanged Low
	G24cd3	zinc finger protein 42 (myeloid-sr		105	185.3347		1.28E+00	4.70E-01	1 Unchanged Low
P41968	N12ef4	melanocortin 3 receptor	225,3987	287	185,5363		-2.81E-01	3.49E-01	1 Unchanged Low
P54920	E03cd4	N-ethylmalelmide-sensitive factor	306,753	267	185.697		-7.24E-01		1 Unchanged Low
Q99487	C19ab8	platelet-activating factor acetylhy		241	185.7091		-2.99E-01	8.00E-02	1 Unchanged Low
Q9H0C8	N15gh7	integrin-linked kinase-associated		109	185.7532	138.517	6.20E-01	-1.49E-01	1 Unchanged Low
P17252	F17ab8	prolein kinase C; alpha	108.3851	74	185.8466	122,7553	7.78E-01	-5.50E-01	1 Unchanged Low
P06493	B01ef5	cell division cycle 2; G1 to S and	108,1696	76.2	185.8754	123.4207	7.81E-01	-5.05E-01	1 Unchanged Low
000139	C14ab7	kinesin heavy chain member 2	140.4141	205	185.9437	177.1988	4.05E-01	5.48E-01	1 Unchanged Low
P54259	N05ef6	dentatorubral-pallidoluysian atroj	165.5295	152	186.0962	167,773	1.69E-01	-1.26E-01	1 Unchanged Low
Q99757	G10cd8	thioredoxin 2	309.0415	163	186,106	219,4587	-7.32E-01	-9.21E-01	1 Unchanged Low
Q9NW68	B12gh2	hypothetical protein FLJ10276	107.4044	242	186.1575	178.6551		1.17E+00	1 Unchanged Low
P16106	O16gh6	H3 histone family; member J	137.9914	135	186,2589	152,9203	4.33E-01	-3.69E-02	1 Unchanged Low
P40692	M06ef6	mult. homolog 1; coton cancer; n		108	186.2872	131.0441	9.15E-01	1,30E-01	1 Unchanged Low
Q99798	E01ab2	aconitase 2; mitochondrial	615,211	291	186.5524		-1.72E+00		1 Unchanged Low
P53618	A11ef3	coatomer protein complex; subur		102	186.6268		-3.54E-01		1 Unchanged Low
Q9UJ69	N24gh2	Toll-Interacting protein	211.4055	259	186.6408		-1.80E-01		
Q9NRM2	P22ef8	zinc finger protein 277	232.9567	227	186.8854				1 Unchanged Low
000070	B06cd6						-3.18E-01		1 Unchanged Low
Q92879 Q75399	H22gh1	CUG triplet repeat; RNA binding		198			-2.16E-01		1 Unchanged Low
Q9BZE4	117cd8	deformed epidermal autoregulate		266			-1.95E-02		1 Unchanged Low
Q9NXB2	G08gh2	G protein-binding protein CRFG	151.2547	214	187.3094	184,3092	3.08E-01	5.03E-01	1 Unchanged Low
		hypothetical protein FLJ20343	259.0672	285	187.4246		-4.67E-01	1.38E-01	1 Unchanged Low
095424	M06ef4	MYLE protein	103.2077	184	187,5574	158,2861	8.62E-01	B.35E-01	1 Unchanged Low
P54577	D14cd3	tyrosyl-tRNA synthelase	145.5748	108	187.5756	146,9084		-4.36E-01	1 Unchanged Low
043692	N03ef7	prolease inhibitor 15	102.5299	163	187.7565	150.9471		6.65E-01	1 Unchanged Low
O60507	J03cd3	tyrosylprotein sulfotransferase 1	190.271	136	188.0843	171.4862	-1.67E-02	-4.83E-01	1 Unchanged Low
Q9BYS9	B15gh5	MMS19-like (MET18 hornolog; S.		117	188.1006		9.22E-01		1 Unchanged Low
P32780	G11ab6	general transcription factor IIH; p		180	188.3175		-4.59E-01		1 Unchanged Low
Q14004	D07ef5	cell division cycle 2-like 5 (cholin		158	188.3656		1.02E+00		1 Unchanged Low
Q15910	L08ab5	enhancer of zeste homolog 2 (Dr		306	188.8314		-1.75E-01		1 Unchanged Low
P47974	H22ab4	zinc finger protein 36; C3H type-l	187.7102	163	188.8801	179.8162		-2.05E-01	1 Unchanged Low
Q13177	E20ab8	p21 (CDKN1A)-activated kinase:	367,7706	232	188,9048		-9.61E-01		1 Unchanged Low
	P16gh8	baculoviral IAP repeal-containing		126	189.0052		-4.11E-01		1 Unchanged Low
P98172	J01ef7	ephrin-B1	122.8828	109	189.5198	140.4039			
Q9BQB1	N04gh6	hypothetical protein FLJ10342	113.906		189.7648				1 Unchanged Low
075472	H11cd5	DnaJ (Hsp40) homolog; subfamil		204	189.9354	158,4309			1 Unchanged Low
075608	K06cd6	lysophospholipase I		374			-2.58E-01		1 Unchanged Low
000193	E14ef3	small acidic protein	357.6112	284			-9.11E-01		
555.50		acidic protein	223.9793	204	190.2397	232.1223	-2.36E-01	3.42E-01	1 Unchanged Low

Q14452	808ab5	glucocorticold receptor DNA blnc	156.8451	147	190.439	164,6255	2.80E-01	-9.75E-02	1	Unchanged Low
P55268	P18ef6	laminin; beta 2 (laminin S)	125.0717		190.5787	155.8763	6.08E-01	2.81E-01		Unchanged Low
Q9P006	A20ef8	milochondrial ribosomal protein t	114.6952	151	190.6578	152.156	7.33E-01	3.98E-01		Unchanged Low
	P07ef1	CGI-111 protein	158,6983		190.6804	175.2567	2.65E-01	1.52E-01		Unchanged Low
	A10gh4 O14gh2	chromosome 11 open reading fra hypothetical protein FLJ20534	70.02507	161 110	190.6944 191.0561	165,3815 123,5579	4.02E-01 1.45E+00	1.59E-01 6.46E-01	1	Unchanged Low Unchanged Low
	P17ab2	ring finger protein 14	227.9827	247	191.1542		-2.54E-01	1.13E-01		Unchanged Low
Q9UBS4	J06ef2	DnaJ (Hsp40) homolog; subfamil		175	191.2168	182.7428	7.25E-02			Unchanged Low
Q12797	J22ab2	aspartate bela-hydroxylase	101,2251		191.3988	126.3618	9.19E-01			Unchanged Low
Q9H4I9	P10gh8	hypothetical gene supported by /			191.4831	192.7801	1.93E-01	3.89E-01		Unchanged Low
O14907 Q14673	E24ef8 B09gh1	Tax interaction protein 1 Bd-2-associated transcription fac	351.7729 97.16912	403 117	191.7684 191.9499	315.6313 135.533	9.82E-01	1.97E-01 2.74E-01		Unchanged Low Unchanged Low
	P01gh2		567.7076		192.0982		1.56E+00			Unchanged Low
Q9BVH4	G21gh6	hypothetical protein MGC3196	164.8572	169	192.4133	175.4765	2.23E-01	3.72E-02		Unchanged Low
P22897	117ab7	mannose receptor; C type 1	77.30638	100	192.4612	123.4215				Unchanged Low
P19012	L04ab6	keratin 15	146.5632	130	192.6097 192.7984	156,5003		-1.69E-01		Unchanged Low
Q14738 O95152	C16cd1 F04cd4	protein phosphalase 2; regulator vesicle trafficking protein	197,6319	183 196	192.7984	195,3686	-2.19E-01			Unchanged Low Unchanged Low
P78426	A09cd3	NK6 transcription factor homolog	198,5291	353	193.0203		-4.06E-02	8.31E-01		Unchanged Low
Q14790	P14ab3	caspase 8; apoptosis-related cys	124,8603	284	193.1013	200.6718	6.29E-01	1.19E+00		Unchanged Low
P07711	N13ab4	cathepsin L	137.16	121	193.1454	150.5861		-1.75E-01		Unchanged Low
P32942	B02ab6	intercellular adhesion molecule 2		140	193,1656	157.6897		-8,57E-03 -5,29E-01		Unchanged Low Unchanged Low
Q9NWS0	L18ab2 F13gh2	ATPase; Na+/K+ transporting; at hypothetical protein FLJ20643	188,9371 81,29108	131 107	194.2503 194.4652	171.3764 127.6547	1.26E+00	3.99E-01		Unchanged Low
Q9NWZ5	006gh2	uridine kinase-like 1	176,188	358	194,4782	242.9739				Unchanged Low
075792	D15cd6	ribonuclease HI; large subunit	100,3515	99.6	194.705	131.5441		-1.12E-02	1	Unchanged Low
P05141	104ab2	solute carrier family 25 (mitochor	454,9832	311	194.7577	320.3918				Unchanged Low
Q9NX38	K22gh2	hypothetical protein FU20457	140,7383	161	195.2668	165.7012		1.95E-01 4.86E-01		Unchanged Low
Q9Y682 P78381	A18cd8 C02cd4	sec22 homolog solute carrier family 35 (UDP-gal	193.7656 276.1892	271 207	195.3918 195.6585	220.1941		-4.14E-01		Unchanged Low Unchanged Low
Q9UMP5	B23ef3	replication initiation region protei		179	195.6957			-5.97E-01		Unchanged Low
Q9P0Q0	C15ef3	hypothetical protein HSPC228	129,6054	241	195.7376	188.838	5.95E-01	8.96E-01		Unchanged Low
Q9UBQ3	M12cd7	polyamine-modulated factor 1	189.4397	172	195.7465	185.7473		-1.39E-01		Unchanged Low
P27540	D22ab2	aryl hydrocarbon receptor nuclea	239.4814	258	195.776	231.071	-2.91E-01	1.07E-01		Unchanged Low
Q9Y2W1 P82267	F03cd5 E15ab4	thyroid hormone receptor-associal adaptor-related protein complex	160,2008 198,6022	248 231	195.7775 195.7793	201.3236	2.89E-01	6,30E-01 2,20E-01	1	Unchanged Low Unchanged Low
P00918	C16ab3	carbonic anhydrase II	283.4506	172	195,9099	216.9569				Unchanged Low
Q9Y663	E07cd8	heparan sulfate (glucosamine) 3-	259.052	99.8	196.1045			-1.38E+00		Unchanged Low
Q9NX40	K18gh2	ovarian carcinoma immunoreacti	119.6155	247	196.1836	187.7106		1.05E+00	1	
P50222	L09ab7	mesenchyme homeo box 2 (grow	165,3048	223	196.5502	195.0839	2.50E-01	4,34E-01	1	
Q13610 P49591	C16cd7 B12cd2	nuclear phosphoprotein similar k seryl-IRNA synthelase	160.8948 152.5268	331 69.6	196.7055 196.7944	229.4375 139.6467		1.04E+00 -1.13E+00		Unchanged Low Unchanged Low
P04554	C19gh5	protamine 3	314.8934	296	196.9835	269.3189		-8.89E-02	1	
P02538	N12ef6	keratin 6A	227.8925	277	197.1759		-2.09E-01		1	Unchanged Low
Q13637	G23cd7	RAB32; member RAS oncogene	392,0158	269	197.2609			-5.43E-01		Unchanged Low
P39086	J02ab3	glutamate receptor; ionotropic; ka		210	197.334			1.88E-01 -5.06E-01		Unchanged Low
Q9H8F9 Q99460	H20gh5 L13ab8	hypothetical protein 24432 proteasome (prosome; macropali	287.08	202 228	197.402 197.4025			-3.89E-01		Unchanged Low Unchanged Low
Q9UKP6	D20gh1	G protein-coupled receptor 14	213,3717	142	197.5131			-5.86E-01		Unchanged Low
Q16589	A04ef5	cyclin G2	99.61434	214	197.5658	170.3526		1.10E+00		Unchanged Low
095316	M03cd5	ribosomal protein S6 kinase; 90k		145	197.5662	162.324		-2.40E-03		Unchanged Low
014545	A09cd7	FLN29 gene product	238,4292	288	197.8936		-2.69E-01			Unchanged Low
Q9Y249 Q92794	P03cd6 K12cd4	homologous to yeast nitrogen pe zinc finger protein 220	191,1146 134,123	224 179	198,1049 198,312	204.4863 170.3523	5.18E-02 5.64E-01	2.31E-01 4.13E-01		Unchanged Low Unchanged Low
P36894	A12ab3	bone morphogenetic protein rece		236	198.4963	194.5315	4.11E-01			Unchanged Low
P79525	L21ab7	MHC class I polypeptide-related	120.8289	125	198.5782		7.17E-01			Unchanged Low
O14947	K04ab6	laminin; beta 3 (nicein (125kD); k		284	199.2636			-3.47E-01		Unchanged Low
Q12899	K16cd3	tripartite molif-containing 26	188.6935	250	199.2998	212.5994	7.89E-02			Unchanged Low
Q9UU5	M24gh8 A18ef2	zinc finger protein 289; ID1 regul zinc finger; DHHC domain contai		137 452	199,4969 199,696	142.4645	1.13E+00 -1.13E+00			I Unchanged Low I Unchanged Low
P46821	J11ab7	microtubule-associated protein 1		136	199,8286		7.21E-01			Unchanged Low
O60906	P04cd1	sphingomyelin phosphodlesteras		2B1	199.835			-7.08E-03		Unchanged Low
P04040	O18ab5	catalase	160.7098	239	199.9283			5.71E-01		Unchanged Low
060709	B16cd5	dynamin 1-like	85,33527	220	199.9492			1.37E+00		Unchanged Low Unchanged Low
Q14714 P39687	A02cd4 F01cd4	sarcospan (Kras oncogene-asso- acidic (leucine-rich) nuclear phos		148 108	200.1413			-1,36E+00 -1,07E+00		Unchanged Low
075341	K16cd4	BRCA1 associated protein	151.895	209	200.166					Unchanged Low
Q9NUN1	D16gh3	hypothetical protein FLJ11259	152.5767	196	200.5435					Unchanged Low
Q9UHR5	L22ef7	transcriptional regulator protein	225,1172	345	200.7483	256.9591	-1.65E-01	6.16E-01		1 Unchanged Low
Q9UMX3	D15gh8	BCL2-related ovarian killer	185.2616	176				-7.26E-02		Unchanged Low
Q02297	J05ef6	neuregulin 1	89,51288	234	201.3025			1.39E+00		Unchanged Low
Q9BQB3 O43491	A21gh6 C22ab4	hypothetical protein MGC5178 erythrocyte membrane protein ba	302.7768	398 144	201.3498		-5.89E-01	3.95E-01 -8.84E-01		1 Unchanged Low 1 Unchanged Low
Q92922	N16cd1	SWI/SNF related; matrix associa		195	201.571			-3.64E-01		1 Unchanged Low
Q16099	J11ef3	glutamate receptor; lonotropic; ka		288	201.66	229,9201	1.17E-02	5.26E-01		1 Unchanged Low
O9532B	B06gh4	hypothetical protein 628	253.2751	240				-7.56E-02		1 Unchanged Low
Q06265	C15cd1	polymyositis/scleroderma autoan		182				1.25E-01		1 Unchanged Low
Q01780 Q9NPF9	K12gh6 G07ef3	 polymyositis/scleroderma autoan HeLa cyclin-dependent kinase 2 		270 170		223.1866		4.57E-01 -5.70E-01		1 Unchanged Low 1 Unchanged Low
Q9P1U0	B03ef4	zinc ribbon domain containing; 1		151				4.48E-01		1 Unchanged Low
Q13480	C02ab5	GRB2-associated binding protein		150		174.7367		-1.98E-01		1 Unchanged Low
095205	O05gh1	C3H-type zinc finger protein; sim	136.7838	267			5.72E-01	9.62E-01		1 Unchanged Low
076091	iticd1	nitrilase 1	194.24	179				-1.17E-01		1 Unchanged Low
Q9Y364	F21ef2 114cd5	DKFZP434J154 protein	208.15	322 213				6.28E-01 2 -2.12E-02		1 Unchanged Low 1 Unchanged Low
P51116 P49915	H19cd4	fragile X mental retardation; auto guanine monphosphate syntheta						-6.13E-01		1 Unchanged Low
		5 Thomps to opinion of the lead	,, _,,				2,,040			

Q9UHR3	O04cd8	nasopharyngeal carcinoma susc	120 0173	198	202 0242	173.3257 7.64E-0	7.005.04	4 11
Q15007	O10cd5	Wilms' tumour 1-associating prof	297 1077	258				1 Unchanged Low
P24941	B03ef5	cyclin-dependent klnase 2	98,60804					1 Unchanged Low
P53634	L01ef7	cathepsin C		214		172.2468 1.05E+00		1 Unchanged Low
Q13123	K03ef7		97.3314	77.9			-3.22E-01	1 Unchanged Low
Q9NUT6		IX cytokine; down-regulator of Hi		153		155.5636 9.16E-01		1 Unchanged Low
Q9Y279		hypothetical protein FLJ11151	285.3106	565		351.7208 -4.77E-01	9.85E-01	1 Unchanged Low
	B23cd7	ig superfamily protein	356.8489	226		262.6019 -7.99E-01	-6.59E-01	1 Unchanged Low
075319	H07cd3	dual specificity phosphatase 11 (250.2151	205	205.1456	220,2678 -2,87E-01	-2.84E-01	1 Unchanged Low
Q9NYH9		hepatocellular carcinoma-associ	287,2586	283	205,223	258,6032 -4,85E-01	-1,99E-02	1 Unchanged Low
Q9UNF0	M20cd7	protein kinase C and casein kina	438,5503	349	205,53			1 Unchanged Low
O60437	D03ab8	periplakin	443.9882	301		316.8358 -1.11E+00		1 Unchanged Low
Q9NZN8	N22cd8	CCR4-NOT transcription complex	189.665	224		206.5153 1.17E-01		
Q14232	A18ab4	eukaryotic translation initiation fa		210		229.9659 -4.12E-01		
O00142	C03ef5	thymidine kinase 2; mitochondria		147				1 Unchanged Low
O60443	N15ab5	deafness; autosomal dominant 5		264				1 Unchanged Low
O95406	J20cd5	comichon-like					1.11E+00	1 Unchanged Low
Q12841	E24cd7	follistatin-like 1	181.7181	138			-3.95E-01	1 Unchanged Low
015217	B04ab5		80.93058	97.7		128.3877 1,35E+00		1 Unchanged Low
P10645		glutathione S-transferase A4	78.38055	91.3		125.424 1.40E+00	2.20E-01	1 Unchanged Low
Q9HA66	E01ab4	chromogranin A (parathyroid sec		283		201,8922 8,38E-01	1.29E+00	1 Unchanged Low
	D09gh5	G protein beta subunit-like	198.1992	192			-4.27E-02	1 Unchanged Low
Q9Y439	B06efB	PRP31 pre-mRNA processing far	175.7234	310	207.0904	230.9065 2,37E-01	8.19E-01	1 Unchanged Low
Q9Y386	G08ef2	CGI-78 protein	325.6509	246	207.7702	259.7061 -6.48E-01	-4.06E-01	1 Unchanged Low
P53990	D21gh1	KIAA0174 gene product	230.1187	401	207.9831	279.676 -1.46E-01		1 Unchanged Low
O95865	Offer3	dimethylarginine dimethylaminoh		157	208.0048	163,3476 7,35E-01		1 Unchanged Low
	B21ef1	immunoglobulin lambda-like poly	142.7367	230	208.1338	193,5142 5,44E-01		1 Unchanged Low
Q15834	117cd7	hepatitis delta antigen-interacting	320.7647	226		251.6031 -6.24E-01		1 Unchanged Low
P18846	H05ab4	activating transcription factor 1	470.9776	206		295.2222 -1.18E+00	-1 195-00	
Q14349	A23cd7	HLA-B associated transcript 8	138,8938	175		174.3142 5.87E-01		
Q9NVM5	M08gh3	hypothetical protein FLJ10637	98.19923	115				1 Unchanged Low
P12955	N18ab7	pepiidase D	133.8157			140.794 1,09E+00		1 Unchanged Low
Q99700	D14cd1	spinocerebellar alaxia 2 (olivopor	101 0044	192 240		178.2362 6.43E-01		1 Unchanged Low
Q9UJA2	E21gh3	chromosome 20 open reading fra	191.0044		209.0555	213.1956 1.30E-01		1 Unchanged Low
Q02246	J21ef6	controlle 20 open reading its		316	209.227	219.8275 6.40E-01		1 Unchanged Low
P48553		contactin 2 (axonal)	108.0043	68.8			-6.50E-01	1 Unchanged Low
	E06cd2	transmembrane protein 1	148.0375	238		198.4177 5.00E-01		1 Unchanged Low
Q99639	K03cd2	suppressor of Ty 5 homolog (S. t	225.1962	296	209.795	243,5051 -1.02E-01	3.92E-01	1 Unchanged Low
Q9NQX5	N02gh4	neural proliferation; differentiation	188,0695	206	209.8245	201,225 1,58E-01	1.30E-01	1 Unchanged Low
Q99799	O08cd6	nuclear RNA export factor 1	205.4132	175	209,9237	196.7655 3.13E-02	-2.32E-01	1 Unchanged Low
Q98ZV1	J05gh7	UBX domain-containing 1	127.7069	236	209.9733	191.2074 7.17E-01		1 Unchanged Low
Q03113	M19ab5	guanine nucleolide binding prote	140,2412	112	209.9975		-3.22E-01	1 Unchanged Low
P01130	B21ab6	low density lipoprotein receptor (301.1179	394	210,369	301.7776 -5.17E-01		1 Unchanged Low
Q9UHN6	L11cd8	transmembrane protein 2	182,3012	130			-4.89E-01	1 Unchanged Low
Q99738	A03cd3	pinin; desmosome associated pri		204	210.6944		2.10E-02	1 Unchanged Low
Q9Y576	N03ef2	ankyrin repeat and SOCS box-cc	121.1139	76.2			-6.69E-01	1 Unchanged Low
P02304	B03gh6	H4 histone; family 2	81.15302	191	210.8651			
P01118	108ab7	v-Ki-ras2 Kirsten rat sarcoma 2 v		239	211.0234			1 Unchanged Low
P15924	K13ef7	desmoplakin (DPI; DPII)	249.9083	263	211.0334			1 Unchanged Low
O43597	A19cd6	sprouty homolog 2 (Drosophila)	119,5051			241.3819 -2.44E-01	7.48E-02	1 Unchanged Low
P09622	106ab3	dihydrolipoamide dehydrogenase		155	211.0573	161.8627 8.21E-01	3.75E-01	1 Unchanged Low
Q08257	G16ef7	composition and designing	546.3029	232	211.1679	263,2128 -7.14E-01	-5.77E-01	 Unchanged Low
Q9UHY8	J09cd7	crystallin; zeta (quinone reductas		301	211,8448	342,4839 -1.28E+00		1 Unchanged Low
P10646	P23cd2	fasciculation and elongation prob	86.8735	95.1	211.8371	131,2705 1,29E+00		1 Unchanged Low
Q92685		tissue factor pathway inhibitor (iii	596,8869	458	212.0146	422.403 -1.49E+00	-3.81E-01	1 Unchanged Low
	L08cd5	Not56 (D. melanogaster)-like pro		217		208.1394 1.21E-01	1.53E-01	1 Unchanged Low
P34949	113ab7	mannose phosphate isomerase	102.2281	135	212.3424	149.8251 1.05E+00	4.00E-01	1 Unchanged Low
Q9NPE9	M21ef8	HSPC055 protein	91.56792	157	212.3544	153.7649 1,21E+00	7.81E-01	1 Unchanged Low
Q99853	J24ef7	forkhead box B1	141.4272	105	212.4773	153.0918 5.87E-01	-4.25E-01	1 Unchanged Low
Q9HD41	C24gh4	CHMP1.5 protein	280.2988	230	212.8608	241.047 -3.97E-01		1 Unchanged Low
Q16625	A10ab8	occludin	859.5688	338	212.9222	470.2634 -2.01E+00		1 Unchanged Low
P45973	L18ef6	chromobox homolog 5 (HP1 alph	164.9514	124	213,1315		-4.13E-01	1 Unchanged Low
Q9NVI9	O12gh3	hypothetical protein FLJ10707	136.0971	136		161.8885 6.47E-01	3.42E-03	1 Unchanged Low
P37173	O09ef7	transforming growth factor; bela i		208		178,0564 9,24E-01	8.90E-01	
Q9UKF7	O21cd8	retinal degeneration B bela	333.6061	287		277.9863 -6.44E-01		1 Unchanged Low
P16870	J23ef7	carboxypeptidase E	81.85341	137		144.0377 1.38E+00		1 Unchanged Low
P00374	D16ab3	dihydrofolate reductase	148.8142	59.6	213.6217			1 Unchanged Low
P19957	M04ab8		202.9357	582			-1.32E+00	1 Unchanged Low
075962	J24cd2		233,7849	197		333.0661 7.60E-02		1 Unchanged Low
P16050	C24ab2	arachidonate 15-lipoxygenase	288.5504			215.0154 -1.28E-01		1 Unchanged Low
	N14cd7	pilin-like transcription factor				196.6194 -4.31E-01		 Unchanged Low
	J16ab6	Participant ideas.	200.9397		214.0069	213,0068 9.09E-02	1.57E-01	1 Unchanged Low
	N15ef1	karyopherin alpha 3 (Importin alp		313	214.0819	295.4165 -7.45E-01	-1.96E-01	1 Unchanged Low
		acid cluster protein 33	222.816	223	214.2958	220.0079 -5.62E-02	6.21E-04	1 Unchanged Low
	G10cd6		219.0427	26B	214.3846	233.924 -3.10E-02	2.93E-01	1 Unchanged Low
	O05ab7	nucleotide binding protein 1 (Min		223	214.5051	236.5867 -3.45E-01		1 Unchanged Low
	K10ef4	AD-003 protein	120.4834	147		160.6424 8.34E-01		1 Unchanged Low
	O20ab8		517.7448			345.2493 -1.27E+00		1 Unchanged Low
	K01ef6	Interferon regulatory factor 1	310.699			306.8604 -5.32E-01		1 Unchanged Low
	A03ab3	ATPase; H+ transporting; lysosor	299.07		215.0012	383.914 -4.76E-01	1.09E+00	
	G19ab6	holocytochrome c synthase (cyto				293.8906 -2.28E-01	7 10F-01	1 Unchanged Low
	C18ef3		103.4737	123	215.2492	147.2535 1.06E+00	2505.01	1 Unchanged Low
	D23ef8		332.5834	276	215 4455	274.7657 -6.26E-01	2.500-01	1 Unchanged Low
	B09ef4	NAD(P) dependent steroid dehyc	253.061	171				1 Unchanged Low
	106cd1	sarcoglycan; alpha (50kD dystro)	358.221			213.0647 -2.31E-01		1 Unchanged Low
	L05ef2	map kinase phosphatase-like pro			215.8058	234.3861 -7.31E-01	-1.47E+00	1 Unchanged Low
	P17gh5	bynothetical eretain 51 193465			215.9987	272.4953 -3.48E-01	2.48E-01	1 Unchanged Low
	E02cd1		204.1242		216.1178	250.8425 8.24E-02	7.03E-01	1 Unchanged Low
			419.7056		216.1184	310.8344 -9.58E-01	-5.00E-01	1 Unchanged Low
			240.4802	235	216.1338	230.6008 -1.54E-01	-3,21E-02	1 Unchanged Low
	P21cd6		188.9561	242	216.7243	215.7355 1.98E-01	3.54E-01	1 Unchanged Low
Q92664	F07ab4	general transcription factor IIIA	318.1862	193	216.7575	242.5829 -5.54E-01	-7.23E-01	1 Unchanged Low

	Q9Y223	N03cd5	UDP-N-acelylglucosamine-2-eph	207.311	295	217.1156	239.9617	6.67E-02	5.11E-01	1 Unchanged Low
•	095816	E21ab3	BCL2-associated athanogene 2	84.7224	94.2	217.4827	132.1461		1.53E-01	1 Unchanged Low
	Q9H0K5 Q9BTJ1	A21gh8 C03ef2	hypothetical protein DKFZp434H ribosomal protein S27-like	193.1934 132.1497	324 411	217.6924 218.2702	244.8706	1.72E-01 7.24E-01	7.45E-01 1.64E+00	1 Unchanged Low
	Q9NRR5	J23gh4	chromosome 1 open reading fran		201	218,4149	185.8791	6.60E-01	5.40E-01	1 Unchanged Low 1 Unchanged Low
	Q9Y5L3	A13ab4	ectonucleoside triphosphate dipir		316	218,4547	256,4569		4.24E-01	1 Unchanged Low
	075351	K06cd5	suppressor of K+ transport defec		311		267.9692		1.78E-01	1 Unchanged Low
	075443	F07cd2	tectorin alpha	183.4591	150	218.7282		2.54E-01		1 Unchanged Low
	Q9H7X4 O00255	K04gh7 K22ab6	KIAA1085 protein	268.2576 134.7209	372 117	218.7527 218.9335	286.2489 156.9374		4.71E-01	1 Unchanged Low
	Q9BV94	D05gh3	multiple endocrine neoplasia I chromosome 20 open reading fra	319,259	316	219.2324			-1.37E-02	1 Unchanged Low 1 Unchanged Low
	014813	D10ab2		226.3067	320	219.2567		-4.57E-02		1 Unchanged Low
	P35269	D15ab4	general transcription factor IIF; p	243.8395	213	219.8975	225.4377	-1.49E-01	-1.98E-01	1 Unchanged Low
	Q12891	E08ef5	hyaluronoglucosaminidase 2	331.3833	311		287.6363			1 Unchanged Low
	O15162 Q14469	L08gh1 O08ab7	phospholipid scramblase 1 halry homolog (Drosophila)	440.6448	597 251	220.6688 221.0242		-9.98E-01 5.48E-01	4.39E-01	1 Unchanged Low
	P06241	P13ab4	FYN encogene related to SRC; F	151.2148	82.5	221.2951		1.06E+00		1 Unchanged Low 1 Unchanged Low
	Q9BVG9	P17gh7	phosphatidylserine synthase 2	225.5812	200	221.5588		-2.60E-02		1 Unchanged Low
	Q9P0P9	K12ef2	hypothetical protein LOC51249	290.4444	434	222.6323	315.5648	-3.84E-01	5.78E-01	1 Unchanged Low
	Q9UMF9	L14cd3	eukaryotic translation initiation fa		242	222,6344	224.2505		2.24E-01	1 Unchanged Low
	Q03527 P31153	L07ab8 J15ab7	proteasome (prosome; macropali		284 134	222,6794 222,7802		-9.88E-01		1 Unchanged Low
	Q04741	D21ef1	methionine adenosyltransferase empty spiracles homolog 1 (Dros		189	222,791		-1.30E-01	-8.75E-01	1 Unchanged Low 1 Unchanged Low
	Q13153	L22ef5	p21/Cdc42/Rac1-activated kinas-	243.198	277	222.8583	247.6258		1.87E-01	1 Unchanged Low
	Q9Y388	H15ef2	CGI-79 protein	100.7214	181	223.2467	168.2598	1.15E+00	8.44E-01	1 Unchanged Low
	Q9HC14	K17gh5	milochondrial ribosomal protein {		200	223.2594	193.9429	4.92E-01	3.31E-01	1 Unchanged Low
	P30281 Q9UFK2	O08ef5 P05ef8	cyclin D3 DKFZP564K1964 protein	107.4859 133.0476	167 210	223.2751		1.05E+00	6.39E-01	1 Unchanged Low
	Q13011	O23ab4	enoyl Coenzyme A hydratase 1; I		273	223.4229 223.7187	188.8934 278 2904		6.60E-01 -3.07E-01	1 Unchanged Low 1 Unchanged Low
	P09601	E20ef7	heme oxygenase (decycling) 1	186.4519	425	223.7447	278.3661	2.63E-01		1 Unchanged Low
	Q9BVZ8	P23gh8	hypothetical protein FLJ14936	126,689	127	224.0692		8.23E-01	6.71E-03	1 Unchanged Low
	Q13889	A11ab5	general transcription factor IIH; p		202	224.1653	179.5227	9.99E-01	8.50E-01	t Unchanged Low
	P49703	P23ab2	ADP-ribosylation factor 4-like	125.6732	91,2	224,1896	147.0234		-4.62E-01	1 Unchanged Low
	Q9GZU7 P10253	G11gh4 M22ab3	nuclear LIM interactor-interacting glucosidase; alpha; acid (Pompe		150 156	224.2596 224.2782			-1.03E+00 -2.69E-01	1 Unchanged Low 1 Unchanged Low
	Q9Y3V5	B18ef8	hypothetical protein DKFZP586F		426	224,4103	318.1897		4.89E-01	1 Unchanged Low
	014773	P17ef7	cerold-lipofuscinosis; neuronal 2:		214			4.04E-01		1 Unchanged Low
	P22061	G13cd1	protein-L-isoasparlate (D-asparta		163	224.5569	168.6387		4.66E-01	1 Unchanged Low
	P14868	K21ab4	aspartyl-IRNA synthetase	117.1946	78.1	225.0996			-5.85E-01	1 Unchanged Low
	Q14938 O75968	D16ef6 P21ef3	nuclear factor I/X (CCAAT-bindin pannexin 1	215.9189	84.9 216		219.072		-1.55E-01 -1.06E-03	1 Unchanged Low 1 Unchanged Low
	Q9H779	C15gh6	hypothetical protein MGC2821	138.4597	313		225.7323		1.18E+00	1 Unchanged Low
	Q9BX72	C02gh8	IFP38	168.9682	172		188,9468	4.18E-01	2.61E-02	1 Unchanged Low
	043563	G24cd1	solute carrier family 22 (organic o		149			7.56E-01	1.53E-01	1 Unchanged Low
	O15142 Q9UJ41	D17ab2 K17ef4	ARP2 actin-related protein 2 hon		215	226,0552			-2.22E-01	1 Unchanged Low
	P05231	F21ef7	putative Rab5 GDP/GTP exchange interleukin 6 (interferon; beta 2)	240.2349	165 673	226.5205 226.529			4.38E-01 1.49E+00	1 Unchanged Low 1 Unchanged Low
	Q9Y2H0	G24ef8	KIAA0964 protein	279.6086	310	226.545	271.9604		1.48E-01	1 Unchanged Low
	Q9UJT9	E15cd8	F-box and leucine-rich repeat pro	287.9128	294	226,762		-3.44E-01	2.98E-02	1 Unchanged Low
	Q9Y306	O23ef2	pelota homolog (Drosophila)	146.8037	190	226.8083	187.7391	6.28E-01	3.69E-01	1 Unchanged Low
	Q9Y6A6 O75300	N04ef3 K18cd2	transcription factor (p38 interactly		171 73.7	226.9539 227.0651	190.8287		-2.97E-02	1 Unchanged Low
	P51798	LO3ef1	ribosome binding protein 1 homo chloride channel 7	278.4282	282	228,319		-2.86E-01	-4.21E-01 1.98E-02	1 Unchanged Low 1 Unchanged Low
	Q14839	C21ab4	chromodomain helicase DNA bin		161	228,322		1.12E+00		1 Unchanged Low
	O50299	O18gh1	KIAA0552 gene product	132.5981	337	228.426	232.6614	7.85E-01		1 Unchanged Low
	Q9Y6F1	H03ab2	ADP-ribosyltransferase (NAD+; ¢		399	228.5355		-1.24E-01	6.80E-01	1 Unchanged Low
	O15212 Q9Y2W2	E04ef3 J14ef2	HLA class II region expressed ge WW domain binding protein 11	250,113 335,0512	292 304	228,5629 228,6314		-1.30E-01	2.22E-01	1 Unchanged Low
	Q9UKN5	O03cd8	PR domain containing 4	252.6208	80.6	228.639			-1.42E-01 -1.65E+00	1 Unchanged Low 1 Unchanged Low
	Q00978	C24cd6	Interferon-stimulated transcription		404			-4.09E-01	4.13E-01	1 Unchanged Low
	Q9UHY0	L08ef1	HP1-BP74	295.9512	267	228.8435			-1.50E-01	1 Unchanged Low
	Q9Y248	L13ef2	HSPC037 protein	92.39515	70,3				-3.95E-01	1 Unchanged Low
	O96019 P10276	E21ab2 K19ab8	BAF53 retinoic acid receptor; alpha	228.9673 250.0312	345 122		267.9206		5.92E-01	1 Unchanged Low
	Q9BUC7	P22gh8	hypothetical protein MGC4172	136.4413	398		254.8265			1 Unchanged Low 1 Unchanged Low
	Q9HAC1	D01gh5	praja t	143.4188	97.5		156.9785	6.81E-01	-5.57E-01	1 Unchanged Low
	Q93078	M18gh6	H2B histone family; member H	179.9656	203				1.72E-01	1 Unchanged Low
	P08572 P15058	N22ef6	collagen; type IV; alpha 2	70.14126	87.3				3.15E-01	1 Unchanged Low
	Q15543	G10ef5 J15cd2	v-rai murine sarcoma viral oncog TAF13 RNA polymerase II; TATA		174 262				-1.59E-01 2.38E-01	1 Unchanged Low 1 Unchanged Low
	O14901	J05cd3	TGFB inducible early growth resi		277				-4.43E-03	
		N10gh8	diphosphate dimethylallyl diphos		257	231.841			-2.87E-01	1 Unchanged Low
	P51965	O13cd3	ubiquilin-conjugating enzyme E2		673				1.40E+00	
	Q9Y3R2	G04cd8	transportin-SR	338.45		232.2301			-6.31E-01	1 Unchanged Low
	O94829 Q92859	G10gh1 M13ab8	importin 13 neogenin homolog 1 (chicken)	160.8646 153.0455					6.17E-01 7.01E-01	1 Unchanged Low 1 Unchanged Low
	Q12846	O02cd2	syntaxin 4A (placental)	323.5547					-1.80E+00	
	P01286	G15ef1	growth hormone releasing hormo	237.6384	263	233.028	244.6261	-2.83E-02	1.47E-01	1 Unchanged Low
	Q9UH92	L01cd8 ·	transcription factor-like 4	172.7135		233.1639		4.33E-01		1 Unchanged Low
	075323 Q9UQ33	P09ab5 P07ef2	glioblastoma amplified sequence microlubule-associated protein; f						-8.78E-02 3.93E-01	
	Q9Y3A7	A17ef2	CGI-101 protein	402.1639		234.2195 234.3057			-9.34E-01	1 Unchanged Low 1 Unchanged Low
	Q9Y2Z0	A17cd7	suppressor of G2 allele of SKP1;		256				4.71E-01	
	P56211	F16ab2	cyclic AMP phosphoprotein; 19 k	276.9734	386	234.3401	299.2676	-2.41E-01	4.81E-01	1 Unchanged Low
	P21127	O13ef1	cell division cycle 2-like 2	159.1108	304				9.32E-01	1 Unchanged Low
	Q07866	B23ab7	kinesin 2 (60-70kD)	254.5932	408	234.5156	299.1796	-1,19E-01	6.82E-01	1 Unchanged Low

Q9BTI3	H11ef8	KIAA1068 protein	228.7025	230	234,5211	231.0835	3.62E-02	8.33E-03		1 Unchanged Low
Q9H2J4 Q9NXE5		hypothetical protein MGC3062	238.7573	256	235.0353	3 243,2412	-2.27E-02			1 Unchanged Low
Q92572		hypothetical protein FLJ20298 adaptor-related protein complex	193.8379	399						1 Unchanged Low
Q9NXT6		FtsJ homolog 3 (E. coll)	254.7485	209 277				1.27E+00		1 Unchanged Low
015321	F05cd6	transmembrane 9 superfamily m		227				1.20E-01 -3.79E-01		1 Unchanged Low
015383	F07cd6	HIV-1 Tal Interactive protein 2;	402.8554	444			-7.73E-01			 Unchanged Low Unchanged Low
O60403 P49354	B06cd8	olfactory receptor; family 10; sul		262			-1.12E-02			1 Unchanged Low
P48039	K24ab4 B07ef6	famesyltransferase; CAAX box; melatonin receptor 1A		193						1 Unchanged Low
043562	C08ab8	solute carrier family 22 (organic	305.0267	208 125				-5.51E-01		1 Unchanged Low
Q9HC89		calpain 10	212.0136	242						Unchanged Low
P00790	M13ef1	pepsinogen 5; group I (pepsinog	256.009	266			-1.13E-01			1 Unchanged Low 1 Unchanged Low
Q9NVP1		DEAD/H (Asp-Glu-Ala-Asp/His)	78.40555	89			1,59E+00			1 Unchanged Low
Q9UBN1 P35659	F23ef3 M22cd3	calcium channel; voltage-depend		360			-1.88E-01			1 Unchanged Low
P54252	N09ef6	DEK oncogene (DNA binding) Machado-Joseph disease (spino	80.02159	77.8				-4.06E-02		1 Unchanged Low
Q14152	L12cd3	eukaryotic translation initiation fa	364.5131	274 268				8.03E-01 -4.43E-01		
Q9Y314	E16ef2	eNOS interacting protein	123.3313	219						1 Unchanged Low 1 Unchanged Low
Q9NRC7 P02533		sirtuin silent mating type informa	1 136.0486	328		233,7238		1.27E+00		Unchanged Low
060907	N02e16 J19cd2	keralin 14 (epidermolysis bullos; transducin (bela)-like 1 X-linked		190				-9.66E-02	1	Unchanged Low
P43155	B16ab3	camiline acetyltransferase	175.9553 288.355	171 190	237.7396 237.8908			-4.19E-02	1	
Q15077	H08ab8	pyrimidinergic receptor P2Y; G-p	389.9746	248				-6.03E-01 -6.52E-01	1	
Q12840	106ab7	kinesin family member 5A	238.3605	282			-7.25E-04		-	
P26599 Q9Y316	A05cd3	polypyrimidine tract binding prote		345	238,4	247.8557	5.71E-01	1.10E+00		Unchanged Low
P24394	M11ef2 O18ab6	C21orf19-like protein interleukin 4 receptor	450,6398	327				-4.62E-01	1	Unchanged Low
P32418	J22gh1	solute carrier family 8 (sodium/ca	309.5199	330 134	238.6201 238.7225		-3.75E-01		1	
P42694	N01ef3	helicase with zinc finger domain	165,7017	146	238.7561			-3.56E-01 -1.79E-01	1	Grid and God Colf
060755	L09cd3	galanin receptor 3	162.6185	268	238.7669		5.54E-01		1	
Q16204 Q12980	A18cd4	DNA segment; single copy; prob		225		218,6477	3.14E-01		1	
P08559	N15cd7 N16ab7	Conserved gene telomeric to alp pyruvate dehydrogenase (lipoam		174	238.9231	198.8702		-7.95E-02	1	Unchanged Low
Q9Y2U9	K02ef3	host cell factor homolog	127.5142	206 154	239.4879 239.623			-8.63E-01	1	
Q9Y2T2	P17cd7	adaptor-related protein complex:	336.9686	395	239.6299		9.10E-01 -4.92E-01		1	
Q13105	K10cd3	zinc finger protein 151 (pHZ-67)	177.9188	243	240.018		4.32E-01		i	
Q9H1J1 Q08752	J12gh5 E01cd1	similar to yeast Upf3; variant A	154.12	183	241.0316		6.45E-01		1	
P38159	F14ab7	peptidylprolyl Isomerase D (cyclo RNA binding motif protein; X chro	342 4602	203 176	241.0389	296,8872	-8.89E-01	-1.13E+00	1	
Q9Y337	A12cd8	kallikrein 5	215.5515	312	241.0429 241.2144	253,2584 256,4089	-5.0/E-01 1.62E-01	-9.58E-01	1	
P25445	A02ef6	tumor necrosis factor receptor su	102.3524	215	241.2257	186,0304		5.36E-01 1.07E+00	1	
Q9BZH1 O43633	P02gh5	membrane-spanning 4-domains;	350.8825	212	241.5154	268.2725		-7.24E-01	i	
Q99536	O12ef4 B03cd6	pulative breast adenocarcinoma vesicle amine transport protein 1		232	241.573		-3.53E-01		1	Unchanged Low
P24557	N11cd1	thromboxane A synthase 1 (plate		400 236	241.6698 241.6841	288.1564		8.44E-01	1	
O75369	G16ab4	filamin B; beta (actin binding prof		459	241.7672	344.1714	-4.56F-01	-4.66E-01 4.69E-01	1	C. C. C. BOL CON
P50453	H10ab8	serine (or cysteine) proleinase in	439.8605	370	242.4005	350.7941			1	
P35680 Q03924	O18cd1 101gh7	transcription factor 2; hepatic; LF		174	242.5943	217,2497	4.28E-02	~4.40E-01	1	
Q9NRQ5	F17gh4	zinc finger protein 117 (HPF9) FN5 protein	257.8571 361,924	301 325	242.7729	267,2646			1	Unchanged Low
Q99867	G12gh7	tubulin; beta polypeptide 4; mem		247	242.8257 243.6773	309.8867 289,1694				
Q9H682	B19gh7	A20-binding inhibitor of NF-kapp:	205.6095	365	243.7949	271,5091		8.28E-01	1	Unchanged Low Unchanged Low
O00148	N12cd5	DEAD/H (Asp-Glu-Ala-Asp/His) t		140		214.3036	-8.71E-02	-8.92E-01	i	Unchanged Low
Q9UG51	F11cd8 N05ef3	TCF3 (E2A) fusion partner (in ch anaphase-promoting complex su		295	244.0153		5.15E-01	7.91E-01	1	Unchanged Low
075175	F12cd8	CCR4-NOT transcription comple:	274.4079	331 253	244.0324 244.0384		-1.69E-01	2.70E-01	1	
Q9GZQ8	C05gh2	microtubule-associated proteins	261.5987	395	244.0725	242.329 300.1702	8.45E-02	1.35E-01 5.94E-01		
P19438	A04ef6	tumor necrosis factor receptor su		197	244.3384	224,9565	6.66E-02		1	
Q9NRR8 O95162	O04gh4 N10ef8	small protein effector 1 of Cdc42	338.2133	329	244.5729		-4.68E-01	-3.97E-02	1	Unchanged Low
095279	J22cd3	peroxisomal short-chain alcohol i polassium channel; subfamily K;		108 127	244,9155	169.1352			1	Unchanged Low
Q9NTW4	N21ef8	DKFZP564A2416 protein	403.4188	219		215,5542 289,1446				
Q9NWM9		hypothetical protein FLJ20730	121.538	207		191.4607			1	Unchanged Low Unchanged Low
P40121 Q92466	P23ab5	capping protein (actin filament); [452.2006	414		370,7982			1	Unchanged Low
O43365	104ab3 K05gh7	damage-specific DNA binding pn		220		188.2839			1	Unchanged Low
Q9NVJ7	O02gh3	homeo box A3 DEAD/H (Asp-Glu-Ala-Asp/His) t	132.7369	196 284	246.2071	191,5647			1	Unchanged Low
Q9NVY5	H08ef7	serologically defined colon cance	182,4598	178	246.7633 246.7633	288,7892 202,4155		-2.44E-01 -3.55E-02	1	Unchanged Low
Q9BY40	G12gh8	CDA11 protein	141.0763		246.8975	173,0132	8.07E-01			Unchanged Low Unchanged Low
Q9NZE2	006ef2	hypothetical protein LOC51321	76.41026	135	247.0025		1.69E+00	8.23E-01		Unchanged Low
P29275 Q9NZE0	K11ab2 P22ef1	adenosine A2b receptor WW domain-containing adapter •	354.5115			264.0659				Unchanged Low
Q03111	N03ab7	myeloid/lymphoid or mixed-linear			247.6094 247.8275		1.58E+00		1	Unchanged Low
Q08499	G21ef6	phosphodiesterase 4D; cAMP-sp			248.1727	211.7891 155.5405	1.39E-01 9.83E-01			Unchanged Low
Q15628	A20ef6	TNFRSF1A-associated via death	214.1641				2.13E-01			Unchanged Low Unchanged Low
Q9Y657 Q9UHQ1	C07cd7 109cd8	spindin	217.9487		248.4185	222.9201	1.89E-01	-1.07E-01		Unchanged Low
P17980	H24ef6	nuclear prelamin A recognition fa proteasome (prosome; macropair				288.0624	-3.32E-01	-4.84E-02	1	Unchanged Low
P45877	119ab8	peptidylprolyl isomerase C (cyclc	165.0716		248.6286 248.9851	257.496 · 194.3167	-4.13E-01			Unchanged Low
O95340	A23cd5	3'-phosphoadenosine 5'-phospho			249.0622	149.5355				Unchanged Low
	F18gh5	hypothetical protein FLJ12448	158.5794	279	249.1478	228.8009	6.52E-01	8.13E-01		Unchanged Low Unchanged Low
O15172 P01298	E11cd4 F05ab8	phosphoserine phosphatase-like				287.4495	-3.68E-01	-1.45E-01	1	Unchanged Low
Q13395	J17cd2	pancreatic polypeptide TAR (HiV) RNA binding protein 1	544.0034 260.7056	194		329.1527 -			1	Unchanged Low
015551	G23ab4	daudin 3	279.9519		249.3777 249.4618	248.0711	-6.41E-02 -1.66E-01			Unchanged Low
					70 10				1	Unchanged Low

P55060	G20ef6	CSE1 chromosome segregation:	322,1588	276	249.5903	282,6907	-3.68E-01	-2.21E-01	1 Unchanged Low
O43286	E13ab3	UDP-Gal:betaGlcNAc beta 1;4- g	316.581	468	249.8072	344.9578	-3.42E-01	5.65E-01	1 Unchanged Low
P04424	J10ab2	argininosuccinate lyase	594,9833	269	250.046	371.2796	-1.25E+00	-1.15E+00	1 Unchanged Low
Q9NWR5	C10ah3	hypothetical protein FLJ20657	171.9643	252	250.1961	224,6496	5.41E-01	5.50E-01	1 Unchanged Low
Q9BUN8	M21gh6	hypothetical protein MGC3067	165.6904	175	250.2847	197.0059	5.95E-01	7.92E-02	1 Unchanged Low
P49760	G01ab4	CDC-like kinase 2	69.23701	97.1	250.3266	138.8925	1.85E+00	4.88E-01	1 Unchanged Low
Q9UEG3	E01gh7	myotubularin related protein 3	367.623	236	250.553	284.5589	-5.53E-01	-6.42E-01	1 Unchanged Low
Q9P0Y6	N07ef1	chromosome 3 open reading fran	268.1158	256	250.6098	258.2636	-9.74E-02	-6.63E-02	1 Unchanged Low
Q92535	M06ab8	phosphatidylinositol glycan; class	218,2502	172	250.9192	213,6071		-3.46E-01	1 Unchanged Low
Q00577	A24gh7	purine-rich element binding prote	839,6	273	251.0214	454.6488	-1.74E+00	-1.62E+00	1 Unchanged Low
075717	L21ab2	AND-1 protein	225.9974	250	251.1522	242.2724	1.52E-01	1.44E-01	1 Unchanged Low
P04278	L09cd1	sex hormone-binding globulin	215.6067	196	251.1944	221.0173	2.20E-01	-1,36E-01	1 Unchanged Low
P51124	G19ab5	granzyme M (lymphocyte met-as-	108,9072	114	251.2498	158.1827	1.21E+00	7.09E-02	
P15259	P04ab7	phosphoglycerate mutase 2 (mus	333.2442	363	251.6606	315.8301	-4.05E-01	1.22E-01	1 Unchanged Low
Q12893	O13cd7	PL6 protein	174.3077	280	252.0241	235.3565	5.32E-01	6.82E-01	1 Unchanged Low
P57738	A11gh2	T-cell leukemia translocation alte	166.5683	232	252.2165	217.0429	5,99E-01	4.80E-01	1 Unchanged Low
P24863	O22ef5	cyclin C	668.8634	523	252.3632	481.4089	-1.41E+00		1 Unchanged Low
Q9Y370	H03ef2	CGI-60 protein	153.6349	157	252.5036	187.603	7.17E-01	2.82E-02	1 Unchanged Low
Q9HD89	E04gh4	found in inflammatory zone 3	189.8748	182	252.7224	208.2163		-6.07E-02	
O14807	N04cd7	muscle RAS oncogene homolog	105.1243	181	252.8147		1.27E+00	7.81E-01	1 Unchanged Low
Q13435	107cd7	splicing factor 3b; subunit 2; 145		268	252.8473	263.6077	-9.65E-02		1 Unchanged Low
Q9UMY4	F19cd8	sorting nexin 12	178.8659	227	253.0136	219.6698	5.00E-01	3.45E-01	1 Unchanged Low
043929	C04ab8	origin recognition complex; subur		354	253.265	291.5955		4.04E-01	1 Unchanged Low
Q9BZQ1	H14gh7	chromosome 1 open reading fran		362	253.5183	278.4192	2.07E-01	7.21E-01	1 Unchanged Low
P55042	K03ef5	Ras-related associated with diab		514	253.5279		-2.46E-01	7.73E-01	1 Unchanged Low
Q92870	K18ef1	amyloid bela (A4) precursor proti		77.9	253.5714		1.74E+00	3.40E-02	
P33778 Q9ULZ3	C19gh7 K22cd8	H2B histone family; member F	169.4431 255.9554	391 517	253.9554 254.0856	271,5522	5.84E-01 -1.06E-02		
Q13485	A20ef5	apoptosis-associated speck-like MAD; mothers against decapents		181	254.0656	199.4935	6.40E-01	1.51E-01	
Q9H684	C24gh5	zinc finger protein 335	326.6826	261	254.3385	280.5396			1 Unchanged Low 1 Unchanged Low
Q9H7F8	O06gh7	hypothetical protein FLJ20958	190.1073	258	254.7196	234,2579	4.22E-01	4.40E-01	1 Unchanged Low
Q92508	B17gh1	KIAA0233 gene product	241.1859	216	255.0491	237.2652	8.06E-02		1 Unchanged Low
Q9UM47	C15ab8	Noich homolog 3 (Drosophila)	64.97745	132	255,0839		1.97E+00		
P18084	E17ef7	Integrin; beta 5	259.2818	129	255.1335	214.6124			
Q99462	K22gh6	ubiquitin-conjugating enzyme E2	207,6691	179	255.2632	213,9424		-2.15E-01	1 Unchanged Low
075619	J10ef7	RNA helicase-related protein	198,1059	335	255.6713	262,7733		7.56E-01	1 Unchanged Low
Q92733	K07cd1	papillary renal cell carcinoma (ira	200,3207	260	256.1407	238.8334	3.55E-01	3.76E-01	1 Unchanged Low
Q15796	H21ab7	MAD; mothers against decapents	376.8422	446	256.2171	359,5423	-5.57E-01	2,42E-01	1 Unchanged Low
P15291	E07ab3	UDP-Gal:belaGicNAc bela 1;4- g	411.9325	403	256,9006		-6.81E-01		
Q99417	107cd8	c-myc binding protein	264.1386	256	256.9088	259.037	-4.00E-02		1 Unchanged Low
P06732	D09ab4	creatine kinase; muscle	202.527	127	257,2272	195,4365		-6.78E-01	1 Unchanged Low
P50570	K13ab5	dynamin 2	248.1389	228	257.2474	244.4335		-1.23E-01	1 Unchanged Low
Q9Y5A9	F20ef2	high-glucose-regulated protein 8	359.385	371	257.3917		-4.82E-01	4.59E-02	
Q92841	B17cd6	DEAD/H (Asp-Glu-Ala-Asp/His) t		270	257.9255	252.4572		2.30E-01	1 Unchanged Low
P51808	B22cd2	t-complex-associated-testis-expn		245	257.9487	228.2619	5.01E-01	4.24E-01	1 Unchanged Low
O95758 P32970	H23cd5 I15ef5	ROD1 regulator of differentiation		462	258.1758		-6.74E-01 -7.79E-01	1.67E-01	1 Unchanged Low
075689	K21cd7	tumor necrosis factor (ligand) su; centaurin; alpha 1	191.7263	456 327	258.4229 258.6362	259,1653	4.32E-01	4.11E-02 7.71E-01	1 Unchanged Low 1 Unchanged Low
P55072	O06cd4	valosin-containing protein	466.7034	363	258.8411		-8.50E-01		1 Unchanged Low
P47813	A16ab4	eukaryolic translation Initiation fa		337	259.0481	360.7943	-9.10E-01		1 Unchanged Low
Q15532	J03cd2	synovial sarcoma translocation; (212	259.1488	220.5805	4.41E-01	1.50E-01	1 Unchanged Low
Q15813	M19cd2	tubulin-specific chaperone e	151.2337	166	259.2747	192.1296	7.78E-01	1.33E-01	1 Unchanged Low
P14373	808cd2	ret finger protein	282.5757	270	259.3599	270.5765			
014980	G02cd3	exportin 1 (CRM1 hornolog; yeas		207	259.7187	235.5264	1.17E-01		1 Unchanged Low
P07942	P20ef6	laminin; beta 1	71,03574	92.4	259.7579	141.0764		3.80E-01	1 Unchanged Low
P53041	C20cd1	protein phosphatase 5; catalytic:	272,5094	195	259.8136	242.7533	-6.88E-02	-4.76E-01	1 Unchanged Low
Q07820	E12ef6	myelold cell leukemia sequence	362.5007	243	260.0329	288,4688	-4.79E-01	-5.78E-01	1 Unchanged Low
P53816	E04cd7	HRAS-like suppressor 3	606.0957	558	260.1565	474.8201	-1.22E+00		1 Unchanged Low
Q9P1C8	E09gh4	hypothetical protein PRO2577	238.9909	205	260.3147	234.8631		-2.19E-01	1 Unchanged Low
P51966	O21cd3	ubiquitin-conjugating enzyme E2		243	260.3185		8.11E-01	7.10E-01	1 Unchanged Low
Q99739	M03cd7	plm-2 oncogene	178.694	274	260.4482	237.5556	5.44E-01	6.14E-01	1 Unchanged Low
P35913	N14ab7	phosphodiesterase 6B; cGMP-sp		173	260.5708	210.8879		-2.07E-01	1 Unchanged Low
P52756	J24cd5	RNA binding motif protein 5	280.6537	230	261.0637	257.2441			1 Unchanged Low
Q9NXX5	K17gh3 006cd5	hypothetical protein FLJ20004	130.163	197			1.01E+00	5.96E-01	1 Unchanged Low
Q14498		RNA-binding region (RNP1; RRA POU domain; class 1; transcription		216			4.16E-01	1.38E-01	1 Unchanged Low
P28069 P51397	A03ab8 E20ab5	death-associated protein	348.1009	206	261.7763	243.064	-4.10E-01	-3.41E-01	1 Unchanged Low
Q9NUI8	C24gh7	pulative methyltransferase	157,3551	312 329	261.9342 262.0287	249.4637		1.06E+00	1 Unchanged Low
000322	M09cd7	uroplakin 1A	223.5286	295	262.0828	260.1175	2.30E-01	3.99E-01	1 Unchanged Low 1 Unchanged Low
P78368	109ab4	casein kinase 1; gamma 2	211.1048	272	262.1137	248.257	3.12E-01	3.63E-01	1 Unchanged Low
Q01081	122cd4	U2(RNU2) small nuclear RNA au		91.3	262.1811	163.878		-5.98E-01	1 Unchanged Low
043193	J15ab5	G protein-coupled receptor 38	69,57316	206	262.301		1.91E+00		
Q9Y3B0	M03ef2	CGI-105 protein	174.5026	177	262.5458	204.6742			
Q9NV80	H05gh3	WD repeat domain 12	178.5139	279	262.6286	239.9609	5.57E-01		1 Unchanged Low
Q9Y473	O18cd4	zinc finger protein 175	141.3794	275	263.1099			9.57E-01	1 Unchanged Low
Q9Y548	1.20gh2	hypothetical protein DJ157A19.1	233.5442	240	263.4412	245.7773	1.74E-01	4.14E-02	1 Unchanged Low
Q13952	N15ef6		241.7469	354	263.6055	286.4173	1.25E-01	5,50E-01	1 Unchanged Low
P18615	K17cd3	RD RNA-binding protein	237.0199	242	263.7642	247.7362		3.25E-02	1 Unchanged Low
P33241	B23ef1	lymphocyte-specific protein 1	94.80267	131			1.48E+00	4.71E-01	1 Unchanged Low
015116	C01ef4	Lsm1 protein	344.8146	479	264.6979		-3.81E-01	4.74E-01	1 Unchanged Low
P50897	A11ab8	palmiloyi-protein thioesterase 1 (142	264.8236		1.06E+00		1 Unchanged Low
0015:5	F20gh8	MADP-1 protein	134.6124	228	264.9564	209.3298	9.77E-01		1 Unchanged Low
Q9Y3A5	G12ef2	CGI-97 protein	469.4567	544	265.0744		-8.25E-01		1 Unchanged Low
Q9BVT1	G03gh6	leukocyte receptor cluster (LRC)	282.5457	328	265.5443		-8.95E-02		1 Unchanged Low
Q9UM44 Q9UGC7	E10cd7	HERV-H LTR-associating 2	180.7238	119	265.5625	188.5666		-5.98E-01 3.22E-01	1 Unchanged Low
#2000/	A05gh3	similar to prokaryotic-type class i	203.3130	362	203.///8	300.0191	-1.26E-01	J.466-01	1 Unchanged Low

P491	146	G01ab8	neuropeptide Y receptor Y2	202 000 4	-						
P546	587	103ab3	branched chain aminotransferas	222.0084	57.1 93.1				-1.94E+00		Unchanged Lov
0608	884	O09cd6	DnaJ (Hsp40) homolog; subfam	330.3137	36				1.41E-01 1.51E-01		Unchanged Lov
Q9N		J01cd7	cofactor required for Sp1 transc	ri 183.0562	15				-2.84E-01		Unchanged Lov
043		G06cd2	tumor protein D62-like 2	283.4636	34				2.95E-01		Unchanged Low Unchanged Low
0758		K22cd2	SCO cytochrome oxidase deficie	156.7184	117				-4.28E-01		Unchanged Low
P494 Q9N		F01gh1	mitochondrial ribosomal protein		35		398.6973		-7.10E-01		Unchanged Low
0154		E15gh2 P18cd2	mitochondrial ribosomal protein	1 473,2712	449				-7.61E-02		Unchanged Low
0148		K04ab5	protein (peptidyl-protyl ds/trans		249						Unchanged Low
0152		106cd6	G protein-coupled receptor 40 ring finger protein 3	222.6002	170				-3.93E-01	1	Unchanged Low
Q9NI		J01ef3	ADP-ribosylation factor GTPase	243.845 · 344.6989	40°		304.1323		7.18E-01		Unchanged Low
0154	191	P22cd5	WD-repeat protein	254.6166	35				-9.76E-01		Unchanged Low
0753		P01ab5	citrate synthase	656.561	312				4.80E-01		Unchanged Low
P570		l12gh3	hypothetical protein FLJ10525	377.5067	349				-1.07E-00	1	
Q9NI		K06gh4	phospholipid scramblase 3	272.0774	334			-2.06E-02	2.96E-01	i	Unchanged Low Unchanged Low
0154		E05cd5	solute carrier family 16 (monoca	r 384.2992	73.5	5 268.3423			-2.39E+00	i	Unchanged Low
Q9BF P103		K09gh6	hypothetical protein MGC3123	127.3745	266		227.363	1.08E+00	1.17E+00	1	
F 100	υ,	P01cd2 A20gh7	related RAS viral (r-ras) oncoger		204				-6.24E-01	1	Unchanged Low
Q9P0)ta	A08ef8	HLA-B associated transcript 4 HSPC142 protein	168.0802	234			6.79E-01	4.79E-01	1	
Q9Y4		L10cd3	insulin receptor substrate 2	196.1899 297.029	346 329						Unchanged Low
Q9UJ	ΣХ	F04ef2	anaphase-promoting complex su		204			-1.35E-01			
Q9H5		001gh6	hypothetical protein MGC2941	333.5834	308			3.64E-01 -3.01E-01			Unchanged Low
Q928		H23cd4	gamma-glutamyl hydrolase (conj		101			1.80E+00			Unchanged Low
Q9UK		K15ab2	ADP-ribosyltransferase (NAD+;		450			-3.10E-01			Unchanged Low Unchanged Low
Q9NV		N13gh2	hypothetical protein FLJ10120	315.7654	301				-7.10E-02		Unchanged Low
0149		107cd2	serine/threonine kinase 6	139.2727	101		170.6789		-4.69E-01	1	Unchanged Low
P5061 Q9H1	-	B11ef5	cyclin-dependent kinase 7 (MO1	286.8535	423		327.4234	-7.56E-02			Unchanged Low
P2185		H08gh7 E13ab4	unc-93 homolog B1 (C. elegans)	348.8713	501		374.1085	-3.57E-01	5.22E-01		Unchanged Low
Q9UG		B14ef2	adaptor-related protein complex	338.7753	373			-3.14E-01			Unchanged Low
P0403	-	D17ab6	protein kinase; AMP-activated; g 3-hydroxy-3-methylglularyl-Coen	425,3015	543			-6.42E-01		1	Unchanged Low
Q9UK		K03gh3	hypothetical protein H41	643.0085	169 272			1.70E-01			Unchanged Low
09547		B18gh1	hypothetical protein YR-29	248.2059	221			-1.23E+00			Unchanged Low
Q9Y31		B16ef8	sentrin/SUMO-specific protease	220.4064	333			1.42E-01 3.15E-01			Unchanged Low
Q9Y3		A20ef2	CGI-36 protein	346.618	338			-3.38E-01			Unchanged Low
Q9285		B09cd6	sema domain; immunoglobulin de	342.6474	183			-3.20E-01			Unchanged Low Unchanged Low
P1610	_	O08gh6	H3 histone family; member C	105.4744	221			1.38E+00		i	Unchanged Low
04314	_	M05ab4	DEAD/H (Asp-Glu-Ala-Asp/His) t	412.7015	349	274,9731	345.6042	-5.86E-01		i	Unchanged Low
Q1354 Q9H6F	-	L12ef6	histone deacetylase 1	375.5725	372	275,2293		-4.48E-01		i	Unchanged Low
Q9NX		D11gh5 M08gh2	hypothetical protein FLJ22313	306.5811	290			-1.55E-01	-8.00E-02	1	Unchanged Low
09476		P03cd5	hypothetical protein FLJ20487 RAN binding protein 9	311.71	394			-1.77E-01	3.38E-01		Unchanged Low
P8290		N16gh8	mitochondrial ribosomal protein (256.3223 261.4618	268			1.06E-01	6.24E-02	1	Unchanged Low
Q9U10		D18ef8	likely ortholog of mouse heat sho	100 R014	343 199			7.73E-02	3.94E-01		Unchanged Low
P2033		M01ef5	RAB3B; member RAS oncogene	214.8304	126				-4.00E-03 -7.65E-01		Unchanged Low
Q0796		B22ab2	Rho GTPase activating protein 1	219.98	218				-1.55E-02		Unchanged Low
P4944		P03cd7	Glutarrate dehydrogenase-2	344.3683	229			-3.18E-01	-5.88E-01	1	Unchanged Low Unchanged Low
Q9NX4		K08gh2	hypothetical protein FLJ20442	493.7568	601				2.83E-01	- 1	Unchanged Low
Q9NV5 00015		J03gh3	chromosome 20 open reading fra		104	277,4945		9.98E-01			Unchanged Low
Q9UI70		L21cd7 B04ef1	brain acyl-CoA hydrolase	173.5669	189		213.4547		1.25E-01		Unchanged Low
P26045		117ab8	nucleolar protein ANKT protein tyrosine phosphalase; no	307,7657	349		311.4328		1.81E-01		Unchanged Low
Q9Y2A		C24ef3	CD209 antigen-like	412,292	315		334,8518				Unchanged Low
Q9BZ8		08gh7	phospholipase A2; group XII	72.21222 175.384	91.3 216			1.94E+00	3.38E-01		Unchanged Low
P35052		24ab4	glypican 1	214.4355	173		223.0558 221.7567	6.64E-01 3.76E-01	2.99E-01		Unchanged Low
Q9H0S		C21gh7	nuclear receptor binding factor-2	80.19669	155	278,4321		1.80E+00	-3.13E-01 9.51E-01		Unchanged Low
Q15654		07ef7	thyroid hormone receptor interac	264.295	238	278,8016	260.4802	7.71E-02			Unchanged Low Unchanged Low
P51948		09ab7	menage a trois 1 (CAK assembly	239.1769	272				1.84E-01	11	Unchanged Low
Q13133		320cd5	nuclear receptor subfamily 1; gro		260	279,1408	272.0232	1.34E-02	-8.70E-02	1 1	Unchanged Low
Q9P0J9 Q15546		122gh3 323cd8	goliath protein	337.3229	215	279,5646	277.1331	-2.71E-01	-6.53E-01	11	Unchanged Low
P28325		09ab5	monocyte to macrophage differer cystatin D		94	279,6415	170.0401	1.03E+00	-5.39E-01		Unchanged Low
O94985		004ef8	calsynlenin 1	371.6792 137.0471	309	279,7092	320.2206	-4.10E-01	-2.65E-01	1 (Unchanged Low
P48634	N	103cd4	HLA-B associated transcript 2	266.8369	205 179	279.921	207.2587 241.9692	1.U3E+00		1 (Unchanged Low
Q9BRL	6 C	15gh7	Splicing factor; arginine/serine-ri-	220,3804	304	280.5497	268.4079		-5.80E-01		Inchanged Low
O94851		06gh1	KIAA0750 gene product	108.6672	172	280,9738	187.2889	3.48E-01	4.65E-01 6.64E-01		Unchanged Low
Q9NW5		20gh3	hypothetical protein FLJ10305	240.4538	314	281.002	278.4551	2.25E-01	3.85E-01		Inchanged Low Inchanged Low
Q9H6X6		19gh5	Cast, interacting molecule	183.3739	302	281.185	255.4932		7.19E-01		Inchanged Low
P41217 P31512		13ab7	antigen identified by monoclonal	325.1642	166	281,3505	257.5072	-2.09E-01	-9.70E-01		Inchanged Low
Q15425		04ab4 105cd2	flavin containing monooxygenase	192.4074	276	281,6555	249.9252	5.50E-01	5.19E-01		Inchanged Low
P40306		05ab8	SA hypertension-associated horr	186.2788	151	281.9205	206.4709	5.98E-01	-3.01E-01		Inchanged Low
Q9NWZ			proteasome (prosome; macropali hypothetical protein FLJ20514			281.9429	400.975	-5.34E-01	3,28E-01	11	Inchanged Low
P57073		22ef8	SRY (sex determining region Y)-I	287.569 202 5886	146 268	281,9498	238.3826			11	Inchanged Low
Q9Y450				230.8266	312	281.9705 282.4288	251.002	4.77E-01	4.06E-01	11	Inchanged Low
P04062	0	08ab3	glucosidase; beta; acid fincludes	295.5318			274.9973 322.2918	2.91E-01	4,34E-01		Inchanged Low
075065		22gn t	phosphodiesterase 4D interacting	168.0201	154	282,6456	201.6823	7 50E 04	3.93E-01 -1.22E-04		Inchanged Low
Q9ULX2		09613	NIMA (never in milosis gene a)-n	288.5578	208	283,1715	259.9871	2 72F-02	-1.22E-01	1 (Inchanged Low
Q99871		06614	urree prime repair exonuclease 2	205.6083	225	283,2504	237.9086		1.29E-01		Inchanged Low Inchanged Low
Q93065		2000	PERBIT family member in MHC.	295 7132	457	283,4269	345.3524 .	6.12E-02	6,28E-01		Inchanged Low
043837 Q9H9L3		12ab7 06gh7	isocitrate dehydrogenase 3 (NAE			283,6422	310.0846	2.99E-01	-2,30E-01		Inchanged Low
Q13740			hypothetical protein FLJ12671	209.5906		283,6912	257.4051	4.37E-01	4.12E-01		Inchanged Low
Q12888			activated leucocyte cell adhesion tumor protein p53 binding protein	231.4355 132.0226	253	283.7582	256,0277	2 045-04	1 205-01	1 0	Inchanged Low
			process	152.5210	223	203,0312	213.3434	1.09E+00	7.48E-01	1 U	inchanged Low

P78318	J13ab6	Immunoglobulin (CD79A) binding	423,8715	512	283.9511	406.4772	-5.78E-01	2.71E-01	1 Unchanged Low
	K24cd1	SRY (sex determining region Y)-I		197	284,0786	275,3736	-2.81E-01	-8.10E-01	1 Unchanged Low
P18065	B19ef1		286,6516		284,2331	284,4952	-1.22E-02	-2.05E-02	1 Unchanged Low
	F22cd8		223.0694	148	284,717	218.587	3.52E-01		1 Unchanged Low
		aldo-keto reductase family 7; me			284.8919		-7.41E-02		1 Unchanged Low
	N03cd7				285,0096		-3.93E-01		1 Unchanged Low
O95169	P14ab8	NADH dehydrogenase (ubiquino			285.0252	154.8111		8.82E-01	1 Unchanged Low
Q99982	C24ef4		63.10609						1 Unchanged Low
Q99470	F14cd2		230,3684		285.1394	277.1416	3.08E-01	4,56E-01	
P16104	N19ab5		234.8351		285.2056	238.3532		-2.68E-01	
Q9UG25	B04ef8	SOCS box-containing WD proteir	213.293		285.5392	298.9147	4.21E-01	9.00E-01	1 Unchanged Low
Q13158	C21cd4	Fas (TNFRSF6)-associated via d	263.0157	281	285.9553	276.784	1.21E-01	9.74E-02	1 Unchanged Low
Q9NW32	D10gh2	hypothetical protein FLJ10346	235.0532	386	286.3524	302.6214	2.85E-01	7.17E-01	1 Unchanged Low
095153	M05cd5	peripheral benzodiazepine recep	155.1824	231	287.1271	224.5106	8.88E-01	5.75E-01	1 Unchanged Low
Q98Q73	H06gh7	hypothetical protein FKSG28	212.0736	297	287.201	265.3817	4.37E-01	4.85E-01	1 Unchanged Low
Q15426	C02ef7	protein tyrosine phosphatase; rei	296.9619	441	287.2583		-4.79E-02		1 Unchanged Low
Q92858	L12ab2	atonal homolog 1 (Drosophila)	611.3779	278	287.2641	392.3029	-1,09E+00	-1.14E+00	1 Unchanged Low
Q16849	P15ab8	protein tyrosine phosphatase; rec		55.6	287.3262	162.9842	9.77E-01	-1.39E+00	1 Unchanged Low
	D07ab7		369,9925	310	287.3935		-3.64E-01	-2.54E-01	1 Unchanged Low
P11279		hypothetical protein MGC14961	103.7251	80.7	287,4071		1.47E+00		1 Unchanged Low
Q96111	116gh7	lagged 1 (Alagille syndrome)	255,0945	347	287.4633	296,4108	1.72E-01	4.43E-01	1 Unchanged Low
P78504	O20ab2			264	287.503	282.0586			1 Unchanged Low
075937	G06ef3		294.9992			194.1988	1.02E+00	1.00E-01	1 Unchanged Low
060575	O16ef4	serine protease inhibitor; Kazal t		152	287.9	257.9759	-5.42E-02		1 Unchanged Low
Q9NYP7	K05gh5	homolog of yeast long chain poly		187	288.1627	_			
095436	H01cd6	solute carrier family 34 (sodium r		425	288.3247	317.3784	2.73E-01	8.34E-01	1 Unchanged Low
O95324	B15ab2	ATP-binding cassette; sub-family	267.704	275	268,4558	277.0901	1.08E-01	3.94E-02	1 Unchanged Low
Q9UQF6	B0Bef3	ribosomal protein L36	519.2003	425	288.5855	410.9115		-2.89E-01	1 Unchanged Low
P13010	K12ef6	X-ray repair complementing defe	174.6996	256	288.9405	239.8403	7.26E-01	5.51E-01	1 Unchanged Low
Q9UJZ1	P03cd8	stomatin (EPB72)-like 2	490.6115	272	289.2003	350,5499	-7.63E-01	-8.52E-01	1 Unchanged Low
	118ab7	matrix metalloproteinase 14 (mer		196	289.8803	204.0657	1.19E+00	6.26E-01	1 Unchanged Low
P50281	808cd1	replication protein A2 (32kD)	100.0172	194	290.3306	194,7022			1 Unchanged Low
P15927			466,4376	454	290.7216				1 Unchanged Low
075496	802ef1	geminin		263	290.7216				1 Unchanged Low
Q9P1E3	C23gh4	presenilins associated rhomboid-						-6,89E-02	1 Unchanged Low
P40189	B14ab6	interleukin 6 signal transducer (g		315	291,3287				
Q9Y259	H03ab4	choline kinase-like	328.1801	375	291.3288		-1.72E-01		1 Unchanged Low
O00633	C12ef5	phosphatase and tensin homolog		189	291.6118				1 Unchanged Low
Q13432	H13cd5	unc-119 homolog (C. elegans)	127.1818	268	291.793			1.08E+00	1 Unchanged Low
Q9Y677	E03ef2	COP9 constitutive photomorphos	188,9367	204	291.9539	227,7917	6.43E-01	1.29E-01	1 Unchanged Low
Q9BW24		KIAA0676 protein	170.3265	240	292,9836	234.5628	7.83E-01	4.97E-01	1 Unchanged Low
Q9NSD9	B12cd5	phenylalanyl-IRNA synthetase be		219	293,111	246.1288	3,71E-01	-5.22E-02	1 Unchanged Low
		hypothetical protein FLJ20695	497,4612	656	293,1504		-7,63E-01	4.00E-01	1 Unchanged Low
Q9NWQ0		hypoxia-inducible factor 1; alpha		276	293.7543		-8.29E-01	-9.18E-01	1 Unchanged Low
QSNWTB				563	294,1533			-3.78E-02	1 Unchanged Low
Q9UPN3		microtubute-actin crosslinking fac		355	294.3016			1.49E+00	1 Unchanged Low
Q96HW0		zinc finger protein 38 (KOX 25)	126.7599					-1.36E-01	1 Unchanged Low
Q14257	F18ab8	reticulocalbin 2; EF-hand caldun		217	295.1469				1 Unchanged Low
Q9NRJ5	J05gh4	poly(A) polymerase beta (testis s		276	295.7341			-4.60E-01	
P23280	P02ab3	carbonic anhydrase VI	137.6155	102	296,1358			-4.28E-01	1 Unchanged Low
Q14677	102gh1	KlAA0171 gene product	330,738	204	296.1674			-6.96E-01	1 Unchanged Low
Q9Y394	H17ef2	CGI-86 protein	471.0368	358	296.7589			-3.97E-01	1 Unchanged Low
Q9UKX5	L18cd7	integrin; alpha 11	99,18133	126	297.0397		1.58E+00		1 Unchanged Low
Q92539	G02gh1	lipin 2	260.7493	235	297,1151	264.3623	3 1.88E-01	l -1.49E-01	1 Unchanged Low
Q9NW01		hypothetical protein FLJ10402	165.6121	135	297.1276	199,1936	8.43E-0	1 -2.97E-01	1 Unchanged Low
QSNUWS		hypothetical protein FLJ11099	228.3252-	247	297,5163	257.4882	3.82E-0	1.11E-01	1 Unchanged Low
Q9BYC4		mitochondrial ribosomal protein \$		408	297,754	394,928	-6,86E-0	1 -2.31E-01	1 Unchanged Low
	F01ab7	lamin A/C	707.7874	294	297.9921		3 -1.25E+00	-1.27E+00	1 Unchanged Low
P02545	A17cd6	serine/arginine repetitive matrix		331			-9.45E-02		1 Unchanged Low
060585		Janus kinase 1 (a protein tyrosin		441			-3.71E-0		1 Unchanged Low
P23458	L12ef5			471			5 -5.44E-0		1 Unchanged Low
Q9BY13	O06gh8	hook3 protein	435.0319	271					1 Unchanged Low
Q9UBC2		epidermal growth factor receptor							
076022	A14cd7	E1B-55kDa-associated protein 5		243					
P33993	K04ef6	MCM7 minichromosome mainten		165					
Q92953	F02cd2	Ric-like; expressed in many tissu		499					1 Unchanged Low
Q9HAF1	P09gh5	hypothetical protein FLJ11730	270.4702	414					1 Unchanged Low
Q9NZ36	P02gh4	uncharacterized hypothalamus p	505,7323	333			5 -7.53E-0		1 Unchanged Low
Q9BQQ3		golgi phosphoproteln 5	208.3671	343			2 5.27E-0		1 Unchanged Low
095214	P09el3	leptin receptor overlapping trans	292.4076	419	300.35		6 3.87E-0		1 Unchanged Low
Q9Y294	A21ef8	DKFZP547E2110 protein	166.5906	243	301.157	3 237.074	9 8.54E-0	1 5,47E-01	1 Unchanged Low
O95810	O01cd4	serum deprivation response (pho		272	301.213	3 284,386	8 1.06E-0	1 -4.11E-02	1 Unchanged Low
P30670	L21ab4	guanine nucleotide binding prote		468	301.889	9 412.818	8 -6.41E-0	1 -1.59E-02	1 Unchanged Low
P51946	O14ef5	cyclin H	180.4218	244					1 Unchanged Low
		retinoblastoma binding protein 6		346					1 Unchanged Low
Q15290	G18cd1	hypothetical protein FLJ20498		190					1 Unchanged Low
Q9NX11			165.3845				6 6.91E-0		1 Unchanged Low
Q9P0S3		hypothetical protein LOC51240	187.2653	229				1 -1.20E-01	1 Unchanged Low
P04554	H11ab8		393,1528	362					
015243		leptin receptor gene-related prof		391			9 -1.22E-0		1 Unchanged Low
043717	G03cd7		284.4848	364			4 9.07E-0		1 Unchanged Low
095876		hypothetical protein LOC51057	172.7128	226			1 8.11E-0		1 Unchanged Low
Q9H8D5			247.105	190	303.483			1 -3.81E-01	1 Unchanged Low
Q9Y3B6		CGI-112 protein	493.6755	28				1 -7.80E-01	1 Unchanged Low
Q9P0S7		hypothetical protein LOC51238	143.0704	139				0 -4.11E-02	1 Unchanged Low
			310.6617	22				2 -4,93E-01	1 Unchanged Low
060630				41:		4 349,751			1 Unchanged Low
095285				22				1 -5.48E-01	1 Unchanged Low
Q08945				25					1 Unchanged Low
043583			203,0088						1 Unchanged Low
095149			186.979	25					
Q9HAS			243,3076	33					1 Unchanged Low
Q9BQ39	9 A23gh6	nucleolar protein GU2	234.3353	16	0 305.273	34 233.259	3.82E-0	01 -5.49E-01	1 Unchanged Low

0153	04 C	06ef6	CD27-binding (Siva) protein	88,75727	153	305.4708	182 3257	1.78E+00	7 935 01	4	
Q153		08gh1	basic leucine-zipper protein BZA		384	305.6252					Unchanged Low
Q9N2		109ef1	CDA14	235.1955	312		345,6465		1.42E-01		Unchanged Low
P260			pentaxin-related gene; rapidly in:				284,2776				Unchanged Low
Q005		24ef5	PCTAIRE protein kinase 1		61.3		154.9189				Unchanged Low
	AWS E			290.1914	359		318.5577		3.09E-01		Unchanged Low
0953			cerebral cell adhesion molecule	52.28353	234	306,6326		2.55E+00			Unchanged Low
0958			Cdc42 effector protein 3	95.65314	208	306.7186	203,4709		1.12E+00	1	Unchanged Low
			arsenate resistance protein ARS:		225	307.4311	240.0311	7.11E-01	2.60E-01	1	Unchanged Low
Q9BX			transmembrane protein induced i		646	308,2116	455.5065	-4.18E-01	6.50E-01	1	Unchanged Low
P367			prolease; serine; 15	390.4608	365	308.5755	354.6324	-3.40E-01	-9.78E-02	1	Unchanged Low
Q9BL			hypothetical protein MGC4614	250,3191	338	308,7431	299.1204	3.03E-01	4.35E-01		Unchanged Low
Q9Y3			Rab acceptor 1.(prenylated)	158,9446	207	308,7948	224.7776	9.58E-01	3.78E-01	1	
Q929	97 D	03ef7	dishevelled; dsh homolog 3 (Drot	240.0941	305	309.2314	284.6254	3.65E-01	3.43E-01		Unchanged Low
, O146		17cd3	ectodermal-neural cortex (with B'	326,0664	189	309.539		-7.50E-02			Unchanged Low
Q9Y2	:55 10	18cd8	px19-like protein	533.9688	455	309,7497	432,9739	-7.86E-01			Unchanged Low
Q166	64 O	17ef4	protein VA**	354,4762	286	310.2736	316.9703				Unchanged Low
P3194	43 0	04ab7	heterogeneous nuclear ribonucle		250	310.5901		-1.15E+00			Unchanged Low
P4984	48 JO		TAF6 RNA polymerase II; TATA	151,1422	133	310.9274		1.04E+00			Unchanged Low
Q049	84 P		heat shock 10kD protein 1 (chap		327	311.1011		-1.20E-01			
P207	19 N	08ef4	homeo box A5	271.4135	326	311.2742	302.8287		2.63E-01		Unchanged Low
Q9H5			hypothetical protein FLJ22795	325,1674	305	311.3753					Unchanged Low
P1902			peplidylglycine alpha-amidaling r		273			-6.25E-02			Unchanged Low
Q925			KIAA0255 gene product			311.913	268.0943	5.06E-01			Unchanged Low
Q129				270.8313	307	312.0297	296.5662	2.04E-01	1.80E-01		Unchanged Low
Q9BV					222	312,5092	282.474	-2.70E-03		1	Unchanged Low
P5336			nudix (nucleoside diphosphate lir		314	312.5575		-1.48E-01		1	Unchanged Low
			partner of RAC1 (arfaptin 2)	293,2566	349	312.9335	318.382	9.37E-02		1	Unchanged Low
P3586			aryl hydrocarbon receptor	249.5731	417	313.1153		3.27E-01	7.40E-01	1	Unchanged Low
Q1314			putative transmembrane protein	490.8698	608	313.616	470.7589	-6.46E-01	3.08E-01	1	Unchanged Low
01474			death-associated protein 6	348.2382	450	313.7398	370.593	-1.51E-01	3.69E-01	. 1	
Q9GZ			wingless-type MMTV integration	362.0025	327	313.9764		-2.05E-01			Unchanged Low
P0817		19eff	decay accelerating factor for corr	705.5781	457	314.0939		-1.17E+00			Unchanged Low
Q9H6			hypothetical protein RG083M05.:	241.9641	291	314.1855	282.3328	3.77E-01	2.65E-01		Unchanged Low
Q9P0			hypothetical protein LOC51235	266.1575	324	314,3982	301.4061		2.82E-01	i	
Q9H2			spinster-like protein	275.9977	394	314.4425	328.1261		5.13E-01		Unchanged Low
Q9NW	VTO D	23gh2	hypothetical protein FLJ20625	420,628	488	314.6523		-4.19E-01			Unchanged Low
Q9Y62	X2 E		protein inhibitor of activated STA		237		233.7882			i	
Q9P1I	P1 F1		done FLB1727 (LOC51215)	419.9659	330	315.0092		-4.15E-01		i	
P2391	9 H		deoxythymidylate kinase (thymid		84.1	315,562		1.35E+00			
P3714	0 D	09ab8	protein phosphatase 1; catalytic :	355 0781	503	315.7751		-1.69E-01	5.03E-01	1	
Q9NV			homolog of rat nadrin	432.3799	670	315.8314				1	Unchanged Low
Q9UM			HIRA interacting protein 5	504.0153	382			-4.53E-01	6.33E-01	. 1	
P1420			folate receptor 2 (fetal)	235,5291	154	315.8386			-4.00E-01		
Q9UH			makorin; ring finger protein; 1	243.752	243	316.0649	235.1893	4.24E-01		1	Unchanged Low
07584			zinc metalioproteinase (STE24 h			316.3297	267.5313	3.76E-01			
0.00			LIM homeobox protein 4	338.221	174	316.3789		-9.63E-02		1	Unchanged Low
P1658				87.11814	84.8	316.6454		1.86E+00		1	Unchanged Low
Q9NV			ADP-ribosylation factor 3	444.3724	348	317.1873		-4.86E-01		1	Unchanged Low
Q9Y3I			hypothetical protein FLJ10439	420.4716	399	317.2936		-4,06E-01	-7.72E-02	1	Unchanged Low
			F-box only protein 7	368.7457	422	317.6515	369.3121		1.93E-01	1	Unchanged Low
Q9H32		21cd8 \		147.0814	348	318.4769		1.11E+00	1.24E+00	1	Unchanged Low
P4990			selenium donor protein	329,031	446	318,9322	364.5082	-4.50E-02	4.37E-01	1	Unchanged Low
00050			serine/threonine kinase 25 (STE	273.696	275	319.3179	289.4017	2.22E-01	7.86E-03		
00027		4ab5	DNA fragmentation factor; 45 kD	328,993	401	319.634	350.0198	-4.16E-02	2.87E-01		
Q0817			splicing factor; arginine/serine-ric	231.1994	193	319.7733	247.9554	4.68E-01	-2.61E-01		
Q9Y3E			small fragment nuclease	120.8094	282	320,0578	240.8904	1.41E+00	1.22E+00		Unchanged Low
Q9P0N		1ef2 r	mitochondrial ribosomal protein t	377.6692	391	320.1382		-2.38E-01	5.03E-02	1	Unchanged Low
Q9BRI		i1gh8 i	hypothetical protein MGC11303:	373.5695	391	320.4318		-2.21E-01	6.60E-02		Unchanged Low
Q9Y31	19 D1	19ef2 (CGI-30 protein	107.1253	123				1.97E-01		Unchanged Low
Q1651	4 M1	17gh 1	TAF12 RNA polymerase II; TATA	116,3618	304		246.9777	1.46E+00			Unchanged Low
P1348	9 D1			224,1663	232		259.1076	5.18E-01	5.00E-02		
Q9998	7 PO			205.9202	194		240,3908	6.42E-01			Unchanged Low
P5618	1 H2			435.0715	319		358.4382				Unchanged Low
P5114	9 N1			329.7976	358		336.6203				Unchanged Low
Q9HD3			CGI-203 protein	257.866	249		276.3612	3.22E-01		-	Unchanged Low
Q9HAM				291.9641	326		313.4126	1.43E-01			Unchanged Low
P0810				535.4871	382						Unchanged Low
O6049							413.3803				Unchanged Low
Q9HBL	3 M1			189.7411	304		272.2579				Unchanged Low
Q9BSJ			* • • • • • • • • • • • • • • • • • • •	165,2938	224 575		237.4707		4.38E-01		Unchanged Low
O6083				549.1645			482.4222		6.53E-02		Unchanged Low
Q9BTM			SAC2 suppressor of actin mutatir		254			5.87E-01		1	Unchanged Low
Q9H4L				371.9412	334	324.5103	343.6429	-1.97E-01	-1.53E-01	1	Unchanged Low
				47.34403	70.6		147.5387			1	Unchanged Low
P50402			merin (Ernery-Dreifuss muscula		304	324.6751	316.6654	1.47E-02	-8.04E-02		Unchanged Low
Q9NZP			alcium binding protein Cab45 pr		239	324.6956	299.844	-4.98E-02			Unchanged Low
P49116		2cd2 r	nuclear receptor subfamily 2; gro	295.9342	224			1.36E-01			Unchanged Low
P11234		4ab8 v	-ral simian leukemia viral oncog	565,699	444	325.5924	444.9874				Unchanged Low
P15408		1ab5 F	OS-like antigen 2	265.6989	437	325.8448		2.94E-01			Unchanged Low
Q9P1Q			AB3A Interacting protein (rabin:	357.8493	253		312.2142				Unchanged Low
P39060				151.6584	212		230.0488	1.11E+00	4.81F-01		Unchanged Low
Q9Y2N		5gh1 k		378.8717			319,0933	-2.13F-01	5.91F-01		Unchanged Low
P02749		2ab2 a	polipoprotein H (bela-2-glycopn			327.1228	320.758	-3.21E-01	9 52E-01		Unchanged Low
Q9P189	9 L20			392.3328			359.0228	-2 62E-01	1 34E-01		Unchanged Low
043822			hromosome 21 open reading fre		297		291.7148	3.01E-01	2 EOF 44		Unchanged Low
Q9UNZ				413.5167		327.9429	420.0654	3 355 04			
Q9Y4G			lynein; cytoplasmic; heavy polyc				376.7971		3.27E-01		Unchanged Low
Q13310			oly(A) binding protein; cytoplasi			328.1257			1.77E-01		Unchanged Low
P45974			biquilin specific protease 5 (iso)	268 1072				5.03E-02			Unchanged Low
		0		200.10/3	310	JZ0.0Z31	304.4284	2.95E-01	Z.39E-01	1	Unchanged Low

P55317	A04ab7	hepatocyte nuclear factor 3; alph		306	328.8562	302.8162			1	Unchanged Low
043364	H06cd8	homeo box A2	511.053	468	329.1292	435,9554	-6.35E-01	-1.28E-01	1	Unchanged Low
P55345	H15ab6	HMT1 hnRNP methyltransferase-	364.8758	307	329.5023	333.8774	-1.47E-01	-2.48E-01	1	Unchanged Low
P48230	O22cd2	transmembrane 4 superfamily me		417	330.825	437.5376				Unchanged Low
Q9BUY3	C21gh5	sorting nexin 6	166,5144	297	330.8949	264,7192	9.91E-01	8.34E-01		Unchanged Low
			275.2503	372	331.0554	326.0947	2.66E-01	4.34E-01		Unchanged Low
000115	B01ef6									
O4331B	M11cd2		186,7739	247	331.1308	254.8269	8.26E-01			Unchanged Low
095572	B08gh1	mitofusin 2	252,7846	277	331.4516	287.175	3.91E-01	1,33E-01		Unchanged Low
P35080	K12ab8	profilin 2	111.4363	85.9	332.2299	176.518	1.58E+00			Unchanged Low
Q9NW29	E24gh3	hypothetical protein FLJ10350	219,5611	173	332.8499	241.9521	6.00E-01	-3.40E-01	1	Unchanged Low
Q9NZE9	O07ef2		219,3916	348	332.8813	299.9861	6.01E-01	6.64E-01	1	Unchanged Low
Q9BYV2		ring finger protein 30	500.2246	338	333.9675	390.6938	-5.83E-01			Unchanged Low
Q9UJX4	F02ef2	anaphase promoting complex sul		269	334.5884	280.1684		1.81E-01		Unchanged Low
000302	B19cd6	calcium homeostasis endoplasmi		382	334.7371	347.7981		2.22E-01		Unchanged Low
	K23cd1			283		302.7027				
Q92886		neurogenin 1	290.2357		335.1054			-3.76E-02		Unchanged Low
014929	P07cd3	histone acetyltransferase 1	344.9153	130	335.4301		-4.02E-02			Unchanged Low
Q9NVK7	M18gh3	chromosome 19 open reading fre		287	335.4383	301,1995	2.54E-01	2,92E-02		Unchanged Low
Q9UBN0	P09ef4	proline rich 3	309,9093	344	335.4526	329,7234	1.14E-01	1.50E-01	1	Unchanged Low
043611	E14cd3	cytoplasmic linker 2	180.9108	220	336.1459	245.5832	8.94E-01	2.B0E-01	1	Unchanged Low
094777	G23cd4	dolichyi-phosphate mannosyltran	294.1356	244	336,7183	291.5641	1.95E-01	-2.71E-01	1	Unchanged Low
Q14669	K05gh1	thyrold hormone receptor interac	486.9829	397	337.0199	406.8782	-5.31E-01	-2.96E-01	1	Unchanged Low
Q9NUR0	B20gh3	hypothetical protein FLJ11196	63,09205	107	337.0769	168,9127	2,42E+00	7,56E-01		Unchanged Low
Q16822	LO8ab8	phosphoenolpyruvale carboxykin		156	337.1655	251.8272		-7.52E-01	1	
	A14cd8			220	337.2906	257.3848	6.48E-01	2.92E-02	i	
Q9Y640		stromal cell derived factor recept								
P48595	C01cd1	serine (or cystelne) proteinase in		410	337,3872	269.7194		2.72E+00		Unchanged Low
P26639	M15cd2	threonyl-IRNA synthetase	355.1309	422	337.8401		-7.20E-02	2.47E-01	1	
P29597	O03cd3	tyrosine kinase 2	209,3427	258	338,1471	268.5377	6,92E-01	3.02E-01	1	•
060671	B02ab8	RAD1 homolog (S. pombe)	188.2716	104	339.1461	210.3983	8.49E-01	-8.59E-01	1	Unchanged Low
Q9NW97	P17gh2	hypothetical protein FLJ10199	262.3124	536	339,3086	379.0815	3.71E-01	1.03E+00	1	Unchanged Low
Q9P027	O23ef8	HSPC133 protein	616,0461	363	339,3102	439,5495	-8.60E-01	-7.62E-01	1	Unchanged Low
095772	E20gh8	hypothetical protein MGC3251	195,9084	227	339.3844	254.0574	7.93E-01	2,12E-01	1	
Q9NR28	N13gh4	second mitochondria-derived act		543	339.8107	384,7351		1.00E+00	i	
P11413	F15ab3	glucose-6-phosphate dehydroger		135	340.0217	244,4906		-9.31E-01	i	
			464.4826	320		375.0149			i	
Q99567	A04ab8	nucleoporin 88kD			340.5878		-4.48E-01			
Q9BZR9	F22gh7	ring finger protein 27	206,7918	398	340.7239	315.1764	7.20E-01		1	
043632	L22cd6	gamma-tubulin complex protein 2		322	341.4805	330,4468		-2.73E-02	1	
Q9Y305	105cd8	Mitochondrial Acyl-CoA Thloeste		475	341.5085	421.5064		8.49E-02	1	Unchanged Low
P17096	L14ef6	high-mobility group (nonhistone o	393.4158	266	341.5301	333.78	-2.04E-01	-5.62E-01	1	Unchanged Low
P28677	E10cd3	visinin-like 1	235,283	241	.341.6123	272.797	5.38E-01	3,76E-02	1	Unchanged Low
Q9UBX5	K04cd6	fibulin 5	79.52593	108	341.7638	176.4964	2.10E+00	4.44E-01	1	Unchanged Low
Q9Y304	E12ef2	ovarian cancer overexpressed 1	326.0044	238	341.8947	301.8585	6.87E-02	-4.56E-01	1	
015379	L16ef6	histone deacetylase 3	198,0005	380	342.3529	306.8418		9.41E-01	1	
Q02790	H16ab5	FK506 binding protein 4 (59kD)	323.5993	240	343,4254	302.2597		-4,33E-01	i	
Q9H240	K12ef5		270,5904	566	343.4339	393.3189		1.06E+00	i	
		Notch homolog 2 (Drosophila)								
076095	P22cd6	jumping translocation breakpoint		227	343,482			-1.56E-01	1	•
Q9Y5Q8	P07cd7	general transcription factor IIIC;		324	343,9958	404.6602		-7.50E-01	1	
Q9NNW5		WD repeat domain 6	254.1607	369	344,0253	322.3381	4.37E-01		1	
P01121	C09ef6	ras homolog gene family; membe		304	344.0732	296,4811	5.09E-01		1	
Q9H0V7	K18gh8	hypothetical protein DKFZp564O	568.248	304	344.2985	405.6684	-7.23E-01	-9.00E-01	1	Unchanged Low
060664	P02cd5	cargo selection protein (mannose	520.0919	232	344,5523	365.5782	-5.94E-01	-1.16E+00	1	Unchanged Low
P56554	Q15cd3	ubiquitin-conjugating enzyme E2	374,4383	417	345,0269	378.959	-1.18E-01	1.57E-01	1	Unchanged Low
000330	O06cd3	Pyruvate dehydrogenase comple		430	345,2213	373.6892	-2.99E-03	3.14E-01	1	Unchanged Low
060784	N17cd5	target of myb1 (chicken)	364,3098	317	345,2431			-2.02E-01	1	
Q9NQZ0	H03ef4	CMP-N-acetylneuraminic acid sy		339	345.3498			-1,22E+00	1	
P52790	A04ab5	hexokinase 3 (white cell)	260,6034	430	345,4621	345.2169		7.21E-01	1	
				277					1	
P52815	B10cd1 O10ef1	mitochondrial ribosomal protein l			345,7776		5,61E-01 1,05E+00			
014983		ATPase; Ca++ transporting; card		223	346.1734				1	
P04259	D01ab7	keratin 6B	534.1837	405	347.6543			-3.99E-01		
P16455	M04ef6	O-6-methylguanine-DNA methylti		307	348,0545				1	
Q9Y5L9	NO4cd6	Snf2-related CBP activator protein		187	348.0925			-6.13E-01	1	
P41214	A16gh1	ligatin	237.0742	307	348,5538			3.72E-01	1	
Q9H6U4	P13gh5	hypothetical protein FLJ21865	419.2747	175	348.8375	314.4039	-2.65E-01	-1.26E+00	1	Unchanged Low
Q15393	L10ef7	splicing factor 3b; subunit 3; 130	242.1115	239	349.6571	277.0735	5,30E-01	-1.59E-02	1	Unchanged Low
P55055	C04gh1	nuclear receptor subfamily 1; gro	325.6291	368	349.8393	347.7942	1.03E-01	1.76E-01	. 1	Unchanged Low
P51805	M03ef4	SEX gene	303.0664	383	350.6414	345.4093	2.10E-01	3.36E-01	1	Unchanged Low
075815	G23ab3	breast cancer anti-estrogen resis		293	350.9571	297.768			1	Unchanged Low
Q9Y2Y9	F19ef2	Kruppel-like factor 13	241.1753	366		319.2905		6.00E-01	1	
Q99633	D04cd3	pre-mRNA processing factor 18	405.046	348				-2.18E-01	-	
P05155	C08ab3			334	351.5074					Unchanged Low
	O16cd7	serine (or cysteine) proteinase in								
Q9UN50		EAP30 subunit of ELL complex	321,9045	326					1	
Q14011	E11ab4	cold inducible RNA binding prote		412	-					Unchanged Low
075807	M02ef3	protein phosphatase 1; regulator		652	352.5048		-7.67E-02			Unchanged Low
Q14457	K19ab3	beclin 1 (coiled-coil; myosin-like	368.4142	678						Unchanged Low
Q02818	J06ef6	nucleobindin 1	330.8747	213			9.39E-02	-6.39E-01		Unchanged Low
015235	L10gh1	mitochondrial ribosomal protein	355.1234	467	353.2438	391,6738	-7.66E-03	3.94E-01	1	Unchanged Low
Q9Y3E0	A02ef2	CGI-141 protein	152.8668	320	353.6986			1.07E+00		1 Unchanged Low
Q9NQZ5		START domain containing 7	277.2941	280				1.19E-02		Unchanged Low
Q9UHW4		protein x 0001	266,9848	291						Unchanged Low
P82914	P04gh7	milochondrial ribosomal protein (417				-9.27E-02		1 Unchanged Low
P25787	L11ef7	proleasome (prosome; macropal		526			-2.86E-01			1 Unchanged Low
P26447	D02cd1			273						1 Unchanged Low
		S100 calcium binding protein A4								1 Unchanged Low
Q9Y605	P02gh8	T-cell activation protein	411.1148	356				-2.10E-01		
Q14776	K14cd4	leucine-zipper-like transcriptiona	202,0365	273			8.15E-01			
manual/										
P30837	C02ab2	aldehyde dehydrogenase 1 famil		451				-4.86E-01		1 Unchanged Low
P09668 Q02313		aldehyde dehydrogenase 1 famil calhepsin H Krueppel-related zinc finger prot	460.8488	451 564 391	356.7735	460.5856	-3.69E-0			1 Unchanged Low 1 Unchanged Low 1 Unchanged Low

040440										
Q13148		TAR DNA blinding protein	322.5816	353	357.1404	344.328	1.47E-01	1.31E-01	1	Unchanged Low
Q16645		FK506 binding protein 1B (12.6)	216.7704	486				1.17E+00		Unchanged Low
	NO2gh8	protein phosphatase 1; regulator	77.52993	71.1				-1.24E-01		Unchanged Low
P57739	F18gh1	claudin 2	188.162	238					i	
O60568	F13cd4	procollagen-lysine; 2-oxoglutarat	332,5306	269				-3.04E-01		
Q9HAU4	N11gh5	E3 ublquilin ligase SMURF2	425.5948	665			-2.51E-01		1	
P49914	J01cd6	5;10-methenylletrahydrofolate sy	340 2344	238						Unchanged Low
O60510	E11cd8	atrophin-1 interacting protein 1;	171 9425	265				-5.17E-01		
Q9BX68		histidine triad nucleotide binding	207.0703	360			1.06E+00		1	
P29084	B02ab5	general transcription factor IIE; p							1	
P09526	C05ef6	PAP18: morehor of PAC	201,3372	225		281.687		-2.17E-01	1	
015471	P01ef4	RAP1B; member of RAS ancage	320.4156	355					1	Unchanged Low
Q9UEE9		leukocyte Immunoglobulin-like re	219.8277	260					1	
Q15011	L03ef3	craniofacial development protein	124.0304	245			1.54E+00	9.79E-01	1	Unchanged Low
	8 A13gh6	homocysteine-inducible; endopla		183			2.94E-02	-9.43E-01	1	
		hypothetical protein MGC2491	170.1162	203		244,6389	1.08E+00	2.57E-01	1	
Q9Y397	A04ef2	zinc finger; DHHC domain contai		495	360.6182	428.8727	-2.58E-01	1.99E-01	1	
Q14596	H17ab7	membrane component; chromoso	303,2476	402	361.5265	355,7499	2.54E-01	4.08E-01	1	on an an agod agon
P23497	C13cd2	nuclear antigen Sp100	331.8298	491	361.9825				i	
Q16633	A24cd1	POU domain; dass 2; associating	103.1231	99.3				-5.39E-02	i	Unchanged Low
Q99961	J12cd1	SH3-domain GRB2-like 1	231.086	247			6.51E-01			
Q07021	N20ab3	complement component 1; q sub-	471 8742	202				-1.22E+00	1	
Q16513	E04cd1	protein kinase C-like 2	546.0111	561	364.4852		-5.83E-01			
Q9H070	H10cd8	adaptor-related protein complex:	353.2901	278					1	
Q13227	A16ab5	G protein pathway suppressor 2	557,5311	352		331,9695		-3.46E-01	1	
P20339	M03ef5	RABSA; member RAS oncogene					-6.12E-01		1	The same of the same
Q9NXX4		hypothetical protein FLJ20005	404.6579	664			-1.46E-01		1	
Q92900	N17cd1	regulator of nonsense transcripts	155.2731	246			1.24E+00		1	Unchanged Low
Q9H2K4	105gh3	thromosome 44 as as a series	328.2926	316				-5.69E-02	1	Unchanged Low
014917		chromosome 11 open reading fre		155	366.4866	240.051	8.83E-01	-3.58E-01	1	Unchanged Low
Q9NZ88	E17ef4 J06gh3	protocadherin 17	250.3859	311	366.6879	309,398	5.50E-01	3.13E-01	1	
		chromosome 6 open reading fran	199.6701	362	367,2681	309,7781	8.79E-01	8.60E-01	1	
Q9BYC8		mitochondrial ribosomal protein t		429	367,9727	351.4981	5.13E-01	7.33E-01	1	
Q15311	C21cd7	ralA binding protein 1	295.4648	411	368,6028	358.3107	3.19E-01	4.76E-01	1	
Q9NWE9		hypothetical protein FLJ10074	455.3747	383	369,4489	402.6031	-3.02E-01	-2.50E-01	1	
095571	G22ef3	protein expressed in thyroid	398.0124	507	369.925	425,0962	-1.06E-01	3.50E-01	1	Unchanged Low
P36941	A09ab7	lympholoxin bela receptor (TNFF	403.0565	454	370.0343		-1.23E-01		1	
P49790	F11cd5	nucleoporin 153kD	264.6175	316	370.2793	317.116	4.85E-01	2.58E-01	i	Unchanged Low
Q99607	C06ab4	E74-like factor 4 (ets domain trar	267.135	268	371,3515		4.75E-01	4.79E-03	i	
Q9UIV1	H05cd8	CCR4-NOT transcription comple:	201.0959	465	371.6771	345.9619		1.21E+00		
Q9Y3E4	O13ef2	CGI-146 protein	107.5317	262	371.8333			1.29E+00	1	
P19367	D17ab3	hexokinase 1	512.0647	215	371.9817			-1.25E+00	1	Unchanged Low
Q14250	J15cd5	enigma (LIM domain protein)	229.1944	218	371.9927	272.9152			1	Unchanged Low
Q9UL33	B20ef2	unknown	343.1533	636	372.4427	450.4535		-7.52E-02		Unchanged Low
Q15019	H03ef5	neural precursor cell expressed:	489.0601	274	372.9279		1.18E-01	8.90E-01		Unchanged Low
Q13084	H05cd6	melanoma-associated antigen re	421.8259	278	373.0512			-8.36E-01	1	Unchanged Low
Q9UMR2	Q18cd7	DEAD/H (Asp-Glu-Ala-Asp/His) t	383.654	423						Unchanged Low
P55855	H06cd2	SMT3 suppressor of mif two 3 ho	454 0014		373.0622	393,1434			1	Unchanged Low
Q13155	G20cd4	JTV1 gene	322.6641	444	373.596			-3.58E-02		Unchanged Low
P78371	H11cd6	chaperonin containing TCP1; sut	322.004 J	409	373.7416	368,4293	2.12E-01	3.42E-01	1	Unchanged Low
P08173	P17ab3	cholinerric recorders manufale	480.3464	430	374.014			-1.60E-01	1	Unchanged Low
060478	E02cd2	cholinergic receptor; muscarinic	444.2461	514	374.4117		-2.47E-01	2.11E-01	1	Unchanged Low
060429	D16gh4	transmembrane 7 superfamily me		438	374.61	362.4989	4.47E-01	6.73E-01	1	Unchanged Low
075935	004cd7	hypothetical protein FLJ12886	142.8322	257	375,0573	258.282	1.39E+00	8.47E-01		Unchanged Low
Q13147		dynaclin 3 (p22)	408.9696	432	375.1224	405.4281	-1.25E-01	7.97E-02		Unchanged Low
Q9NPH4	J05ab2	abl-interactor 2	299.098	249	375.379	307,9098	3,28E-01	-2.63E-01	1	Unchanged Low
Q9UJ96	E14gh3	LUC7-like (S. cerevisiae)	207.38	302	376.3768	295,4166	8.60E-01	5.45E-01	1	Unchanged Low
	C13cd8	potassium voltage-gated channe	582.717	448	376.4373	468.9343	-6.30E-01			Unchanged Low
P17081	C13ef6		293.1604	417	376.4996	362,2816	3.61E-01	5.09E-01		Unchanged Low
P11166	J05ef5	solute carrier family 2 (facilitated	351.7333	393	376.564	373.8343	9.84E-02	1.61E-01		Unchanged Low
Q14820	D12ef6	splicing factor 1	382.846	408	376,7779	389,2167		9.19E-02		
Q92935	106ab5	exostoses (multiple)-like †	659,9005	276	376,8548	437.6869				Unchanged Low
P31749	E20ef6	v-akt murine thymoma viral onco	385.4669	303	376,9169	355,1851	-3 24F-02	-3 46E-01		Unchanged Low
Q13454	L05cd7	Putative prostate cancer tumor si		363	377.0852		2.06E-01	1,50E-01		Unchanged Low
P35520	C24ab3		70.85109	68,1	377,1148	172.0203	2.41E+00			Unchanged Low
P53602	M01ab7	mevalonate (diphospho) decarbo		178	378,114	412.6357	.0 E1E 04	1.045.00		Unchanged Low
Q9H7G3	C04gh6		333.6677	207		206 2067				Unchanged Low
Q13123	C06ab7	IK cytokine; down-regulator of HI		439		306,3867 358,3456	1,81E-01			Unchanged Low
O14744	G14cd6	SKB1 homolog (S. pombe)	348.0026				1.21E-01		1	Unchanged Low
Q9Y566	D18ef8		209.2687		378,4165			2.37E-01		Unchanged Low
Q9BUJ9	A03gh7			281	379.5197	290.0718		4.27E-01	1	Unchanged Low
P51790	102ab5		385.0517	472		412.4352			1	Unchanged Low
O14547	J09cd6		330,2251	237	380,3105	315.7621	2.04E-01			Unchanged Low
Q99753	A05cd7	PRP8 pre-mRNA processing fact		256		291.9777	6.71E-01	1.01E-01	1	Unchanged Low
Q9NQC8			257.1873	397	380,7894	345,0384	5.66E-01	6.27E-01	1	Unchanged Low
043919	H15gh4 M01ef6	hypothetical protein LOC56912	185.2607	276	381.0416	280.7773				Unchanged Low
			206.6586	105	381.2059	230.823	8.83E-01			Unchanged Low
P78345 P54257	F15cd6	ribonuclease P (38kD)	321.791	530	381,5687	411.2575	2.46E-01	7.21E-01		Unchanged Low
	A03ef5	huntinglin-associated protein 1 (r		322	382.6029	384,9948				Unchanged Low
Q9Y3F4	G18cd7	unr-interacting protein	390.9437	335	382.6142	369,3734	-3.11E-02	-2,25E-01		Unchanged Low
P29536	F14cd7		343.3731	209	382.9438		1.57E-01			Unchanged Low
075925	K03cd4	protein inhibitor of activated STA		606		471.2944				Unchanged Low
Q9UMX5	J23el4	secreted protein of unknown func	344.8668	306	384,5676	345,2233	1.57E-01			Unchanged Low
Q9UM29	N02cd7	Interleukin enhancer binding fact	348.7866	278		337.1554	1.42E-01			
Q92905	P03ef6	COP9 constitutive photomorphoc	257.5597		386.1452					Unchanged Low
	L20ef1	G-protein coupled receptor SALF	734.6282	277				6.06E-01		Unchanged Low
Q9P2W3	G01ef3	guanine nucleotide binding prote		81.4		179,6321				Unchanged Low
Q9BV17	N14gh5	mitochondrial ribosomal protein !		520						Unchanged Low
Q9Y4L1	B23cd6	oxygen regulated protein (150kD		372		424.8293	0.395-02	4.996-01		Unchanged Low
Q02363	F18ab7	inhibitor of DNA binding 2; domir		359	388 3553	390,9975	9.202-02	-1,492-01		Unchanged Low
		or or a containing 2, contain		333	JU075255	410.6421	3.20E-01	-4.33E-01	1	Unchanged Low

Q9Y5J8	E06cd8	translocase of inner mitochondria	440.2763	542	388.4554	456.8557		2.99E-01	1 Unchanged Low
	E22ef1	G protein-coupled receptor 26	499.1316	450	389,582		-3.57E-01		1 Unchanged Low
Q99501	N11cd6	growth arrest-specific 2 like 1	359.8065	540	389.6398	429.6515	1,15E-01	5,84E-01	1 Unchanged Low
Q00613	M17ef5	heat shock transcription factor 1	363.3847	375	389,8695		1.01E-01	4.71E-02	1 Unchanged Low
Q9H5Z1	H16gh4	DEAD/H (Asp-Glu-Ala-Asp/His) t		124 115	389,9835		1.51E+00		1 Unchanged Low
P09661 P54709	P24cd1 P05el5	small nuclear ribonucleoprotein r ATPase; Na+/K+ transporting; be		432	390.2085 391.8135		-1.20E-01	-1.94E+00 2.22E-02	1 Unchanged Low 1 Unchanged Low
P13164	M06ef5	Interferon induced transmembrar		256	392,1204			2.50E+00	1 Unchanged Low
Q9Y6E2	A23ef8	HSPC028 protein	329.8857	363	392.2431		2.50E-01	1,37E-01	1 Unchanged Low
Q9NVG2	B13gh3	vacuolar protein sorting 35 (yeas	359.0233	557	392.2574	436.0959	1.28E-01	6,34E-01	1 Unchanged Low
P27448	E10ei5	MAP/microtubule affinity-regulation		369	392,3918		-1.97E-02		1 Unchanged Low
015438	P18cd3	ATP-binding cassette; sub-family		488	392.8159		-6.54E-01		1 Unchanged Low
Q9UL48	104ef2	CK2 Interacting protein 1; HQ002		215	393,4111	261,8209	1.16E+00	2.88E-01	1 Unchanged Low
O94763 Q13190	A11cd4 I19cd2	RPB5-medialing protein syntaxin 5A	333.6215 179.6148	533 235	393,9448 393,985	420.3384 269.5995	2.40E-01 1.13E+00	6.77E-01 3.89E-01	1 Unchanged Low 1 Unchanged Low
Q13122	D21ef3	EBNA-2 co-activator (100kD)	381,3069	375	394.5999	383.7948		-2.22E-02	1 Unchanged Low
075312	LO1cd4	zinc finger protein 259	348.7542	422	394.8016	388.3905	1.79E-01	2.74E-01	1 Unchanged Low
P37837	D20cd2	transaldolase 1	399.2717	326	395,693	373.7275			1 Unchanged Low
075762	G19cd6	ankyrin-like with transmembrane	222.8517	364	395.7427	327.3962	8.28E-01	7.06E-01	1 Unchanged Low
Q99959	K19ef3	retinoic acid receptor responder		257	398.0287	312.0672		-1.44E-01	1 Unchanged Low
Q9UNP2	H15cd2	solute carrier family 1 (neutral an		138	396.5014			-1.15E-01	1 Unchanged Low
P56524 Q9H644	C23cd6 C19gh6	histone deacetylase 4 hypothetical protein MGC5585	328.6037 140.9811	303 228	397.1317 397.2624	342.9334 255.3931	1.49E+00	-1.17E-01 6.93E-01	1 Unchanged Low 1 Unchanged Low
Q13627	O19ab4	dual-specificity tyrosine-(Y)-phos		327	397.3922			-6.06E-01	1 Unchanged Low 1 Unchanged Low
Q03405	001ef5	plasminogen activator; urokinase		396	398.2456			-3.38E-01	1 Unchanged Low
Q9NPQ8	L18gh4	likely ortholog of mouse synembr		443	398.4083	404.6217	9.57E-02	2.48E-01	1 Unchanged Low
Q15375	A10ab5	EphA7	243.0574	290	398,5414	310,569	7.13E-01	2.55E-01	1 Unchanged Low
P05114	G04ab7	high-mobility group (nonhistane (278	398.7327	291.2748	1.02E+00	4.99E-01	1 Unchanged Low
Q9UKU4	C13ef2	retinal short-chain dehydrogenas		497	399.4413	422.4244	1.07E-01	4.23E-01	1 Unchanged Low
P08195 Q15545	E21ab7 J11cd2	solute carrier family 3 (activators TAF7 RNA polymerase II; TATA	442,5453 400,689	345 526	399.5678 401.1308	395,768 442,5426	1.59E-03	-3.58E-01 3.92E-01	1 Unchanged Low 1 Unchanged Low
Q13545	D24gh8	hypothetical protein FLJ14511	246.3293	225		290.8781		-1.31E-01	1 Unchanged Low 1 Unchanged Low
Q05397	L20ef5	PTK2 protein tyrosine kinase 2	389.8718	440		410.2909			1 Unchanged Low
Q9Y6G1	C21ef8	PTD011 protein	434.6906	483	401.5119	439.8467		1.53E-01	1 Unchanged Low
P41212	O23ef5	els variant gene 6 (TEL oncogen		502	401.8428	406.6035	3.48E-01	6.70E-01	1 Unchanged Low
014737	G05cd5	programmed cell death 5	498.729	430	402.3091			-2.13E-01	1 Unchanged Low
Q90H89 Q96D21	A10ef3	signal recognition particle 68kD RASD family; member 2	212.5357 395.909	247	402.3276	287.2236	9,21E-01	2.16E-01	1 Unchanged Low
Q9Y411	C20ef8 J05ef1	myosin VA (heavy polypeptide 1:		486 353	402.4304 402.453	427.9499 375.8247	2.36E-02	2.94E-01 -7.23E-02	1 Unchanged Low 1 Unchanged Low
Q9Y5L4	L14ef7	translocase of inner mitochondria		369	404.0738	354.9111		3.39E-01	1 Unchanged Low
075439	N22cd4	peptidase (mitochondrial process		184	404.5715			-1.48E+00	1 Unchanged Low
P55010	K22ab4	eukaryotic translation initiation fa	427.4989	296	405.4996	376.3076	-7.62E-02	-5.31E-01	1 Unchanged Low
Q9BY44	G10gh8	CDA02 protein	453.6064	401	405,956			-1.78E-01	1 Unchanged Low
Q13164	B21ef5	mitogen-activated protein kinase		256	406,234	290.5304	9.58E-01	2.93E-01	1 Unchanged Low
P53667 Q9UNH6	P08ab6 F05ef2	LIM domain kinase 1 sorting nextn 7	359.4655 479.6898	392 396	406.5183 406.7812	386.0829	1.77E-01	1.26E-01 -2.78E-01	1 Unchanged Low 1 Unchanged Low
O00258	K17cd4	tryptophan rich basic protein	339.3172	500	407.1504	415.5431	2.63E-01	5.60E-01	1 Unchanged Low
Q9Y310	112ef3	hypothetical protein	458.2693	546		470.3717		2.52E-01	1 Unchanged Low
Q9Y368	D13cd8	parvin; beta	114.232	159	407.3882	226.9908	1.83E+00	4.80E-01	1 Unchanged Low
P14866	H13ab6	heterogeneous nuclear ribonucle		274	407.4025			-6.92E-01	1 Unchanged Low
Q13885	K01cd3 J06cd1	tubulin; bela polypepiide	441.2569	413	407.8683			-9.41E-02	1 Unchanged Low
O43765 O14632	B12ef4	small glutamine-rich tetratricoper homeodomain interacting protein		438 535	408.6453 408.8773	408.6939 446,1081	1.09E-01 5.18E-02	2.10E-01 4.40E-01	1 Unchanged Low
P31939	L10ab2	5-aminoimidazole-4-carboxamide		386	409.3069			-3.76E-01	1 Unchanged Low 1 Unchanged Low
P28331	P18ab8	NADH dehydrogenase (ublquinor		417	409.3283			-2.35E-01	1 Unchanged Low
P41970	P05ef6	ELK3; ETS-domain protein (SRF	205.9453	359	410.8037	325,3706	9,96E-01	8.03E-01	1 Unchanged Low
Q92973	J22ab6	karyopherin (importin) beta 2	393.6551	234	410.9425	346.1733		-7.51E-01	1 Unchanged Low
P36507	J10ef5	mitogen-activated protein kinase	395.22	437	411.4052	414.6953	5.79E-02	1.46E-01	1 Unchanged Low
Q16512 O43447	F21ab8 M08cd6	protein kinase C-like 1 peptidyl prolyl isomerase H (cycl-	456.2727 179.441	486 222	411.8245 411.9165		-1.48E-01 1.20E+00	9.18E-02 3.05E-01	1 Unchanged Low
P30040	M13ef3	chromosome 12 open reading fre		296	412.2064	351.5665		-2.29E-01	1 Unchanged Low 1 Unchanged Low
095399	C19cd7	urolensin 2	452.0721	366	413.1451			-3.04E-01	1 Unchanged Low
	M01ab3	UDP-Gal:belaGal beta 1;3-galac		290	413.3454			-7.87E-01	1 Unchanged Low
Q12792	N11ab8	protein tyrosine kinase 9	476.3952	364	413.4208			-3.87E-01	1 Unchanged Low
Q14318	J02cd7	FK506 binding protein 8 (38kD)	461.2322	432				-9.35E-02	1 Unchanged Low
075955 Q9BVYB	N10cd5 B09gh7	flotillin 1 hypothetical protein MGC5499	304.6674 490.2382	310	415.4364			2.68E-02 -2.35E-01	1 Unchanged Low
Q16288	019ef5	neurotrophic tyrosine kinase; rec		417 186	416.02 416.4128			5.53E-01	1 Unchanged Low 1 Unchanged Low
075380	J20ab8	NADH dehydrogenase (ubiquinor		413	416.4836			-1.21E-01	1 Unchanged Low
Q9BY51	B15gh8	hypothetical protein GL009	429.1998	354	417.3575			-2.80E-01	1 Unchanged Low
Q9BVK3	E19gh8	hypothetical protein MGC2463	462.5828	387	417.7046			-2.56E-01	1 Unchanged Low
Q9PON3	A24ef2	hypothetical protein FLJ10597	137.7487	289	417,999			1.07E+00	1 Unchanged Low
P24534	D03ab4	eukaryotic translation elongation		328	418.0152			-8.11E-02	1 Unchanged Low
P49336 P34130	C01ab4 P16cd2	cyclin-dependent kinase 8 neurotrophin 5 (neurotrophin 4/5)	426.3751	281	418,121			-6.00E-01 -5.53E-01	1 Unchanged Low
075425	P06gh5	hypothetical protein AF053356_(339	418.6011 419.3637	323.0643 366.1692		-7.34E-03	1 Unchanged Low 1 Unchanged Low
Q15293	F16ab8	reliculocalbin 1; EF-hand calcium		250	420.5597	304,1243	7.99E-01		1 Unchanged Low
095376	D08ab2	arladne homolog 2 (Drosophila)		322		332.2465			1 Unchanged Low
075489	J16ab8	NADH dehydrogenase (ubiquinor	375.7034	327	422.3772	375.1108	1.69E-01	-1.99E-01	1 Unchanged Low
P28482	B15ef5	mitogen-activated protein kinase		452				-1,28E-01	1 Unchanged Low
P20823 O15250	B09cd1 I04cd4	transcription factor 1; hepatic; LF X-prolyl aminopeptidase (aminop		527				7,09E-01	1 Unchanged Low
P55769	P22ab8	NHP2 non-histone chromosome		542 479				4,57E-02 1,80E+00	1 Unchanged Low 1 Unchanged Low
094817	N21ab2	APG12 autophagy 12-like (S. cer		402	424.3577			-6,52E-01	1 Unchanged Low
075352	K08cd5	mannose-P-dollchol utilization de	272.444	523	424.8266			9.41E-01	1 Unchanged Low
Q16585	116cd1	sarcoglycan; bela (43kD dystropl	256.2113	293	425.1172	324.9166	7.31E-01	1.96E-01	1 Unchanged Low

Q9H7X	7 B16gh5	hypothetical protein FLJ14117	261.9578	463	425.3725	383.5805	6.99E-01	8.23E-01	1.1	Unchanged Low
015173	3 116cd6	progesterone receptor membrane	295.0481	244			5.29E-01	-2.74E-01		Unchanged Low
Q9GZX	(9 A16gh4	twisted gastrutation	84.61732	147	426,1124		2.33E+00	7.99E-01		Unchanged Low
Q16594		TAF9 RNA polymerase II; TATA	171.3507	520			1.32E+00	1,60E+00		Unchanged Low
000479		high-mobility group (nonhistone o	265,695	354	426,7642	348.9168	6.84E-01	4.15E-01		Unchanged Low
P54760		EphB4	528,7066	413	427.0543	456.4179	-3.08E-01			Unchanged Low
P25490		YY1 transcription factor	481.2844	530	427.184		-1.72E-01	1.39E-01		Unchanged Low
Q9UJY		putative nucleotide binding prote		431		454.8441	-2.44E-01	-2.31E-01	1.5	Unchanged Low
Q09666		AHNAK nucleoprotein (desmoyol		399	427.382B	406,041	1.24E-01	2.41E-02	1 1	Unchanged Low
O15305		phosphomannomulase 2	325.0777	480	428.3778		3.98E-01	5.63E-01		Unchanged Low
O60492		7-dehydrocholesterol reductase	388.5228	213				-8.68E-01	1 1	Unchanged Low
Q9UND		nuclear pore complex Interacting	280.8175	312			6.10E-01	1.54E-01		Unchanged Low
Q9HB1		potassium channel; subfamily K;	473.5589	408			-1.43E-01		1 1	Unchanged Low
Q9Y38- Q14107		CGI-110 protein	290.9669	331	429.1497	350.4808	5.61E-01	1.87E-01		Unchanged Low
		D123 gene product	387.0465	405	429,9331	407.3091		6.52E-02		Unchanged Low
P52435 Q9Y262		polymerase (RNA) II (DNA direct		461	429,9488		-2.94E-01			Unchanged Low
P10163		eukaryotic translation initiation fa		311	430.6788		-4.58E-01			Unchanged Low
P38378		proline-rich protein BstNI subfam		251	431.1299		-2.74E-01			Unchanged Low
P48735		protein transport protein SEC61 :		369	431.9958	478.7225	-5.57E-01			Unchanged Low
Q9NPF		isocitrate dehydrogenase 2 (NAE		338	432,1616	456.8577	-4.74E-01			Unchanged Low
014616		DNA methyltransferase 1-associa zinc finger protein-like 1		382			-6.26E-02			Unchanged Low
Q00839		helerogeneous nuclear ribonucle	194,5871	348 410			1.15E+00			Unchanged Low
O95563		DKFZP564B167 protein	390.848	389	432.5188 432.8082	419.7665 404.2972		-2.18E-02		Unchanged Low
Q92974		rho/rac guanine nucleotide excha		124	433.2386			-5.96E-03		Unchanged Low
Q9UEE		serine/threonine kinase 17a (apc		472	433.4221		1.61E+00			Unchanged Low
Q9NXB		BTB (POZ) domain containing 2	175.3887	206	433.4811	434.4129 271.6819	1.24E-01 1.31E+00			Unchanged Low
Q9NXF		O-linked mannose beta1;2-N-ace		353	434.2927		-1.37E-01			Unchanged Low Unchanged Low
P98171		Rho GTPase activating protein 4		231	435,3585	345.0241		-6.77E-01		Unchanged Low
P56192		methionine-tRNA synthetase	263.0076	295	435.894	331.1505	7.29E-01	1.63E-01		Unchanged Low
O95411	K09cd5	TGFB1-induced anti-apoptotic fa		310	436.2684	326.6136	9.02E-01	4,10E-01		Unchanged Low
P09132	E23cd2	signal recognition particle 19kD	227.8193	595	438,5994	419.7603	9.38E-01	1,38E+00		Unchanged Low
Q92537	O24gh1	KIAA0247 gene product	291.1054	232	437,1141	319,914		-3,30E-01		Unchanged Low
P49761	G03ab4	CDC-like kinase 3	458,9286	562	437.5601		-6.88E-02	2.91E-01		Unchanged Low
Q9HDC		junctophilin 3	346,7829	388	439.282	391.3835	3.41E-01	1.62E-01		Unchanged Low
P98160		heparan sulfate proteoglycan 2 ()	505.6962	372	439.9544	439.0897	-2.01E-01	-4.44E-01		Unchanged Low
Q9NQV		PR domain containing 10	369,157	514	440.5347	441.3923	2.55E-01	4.79E-01		Unchanged Low
Q9NRX		protein x 013	324.5306	345	440.8995	370.1383	4.42E-01	8.82E-02		Unchanged Low
O76084		catenin (cadherin-associated pro		211	441.7617	374.6479	-9.29E-02	-1.16E+00		Unchanged Low
P23197	E19cd7	chromobox homolog 1 (HP1 beta		362	441.9595		-1,22E-01		1 (Unchanged Low
Q99439		calponin 2	429.1961	304	442.4148	391,978		-4.96E-01		Unchanged Low
O14656		dysionia 1; torsion (autosomal do		533	442.9429	463.6417	9.27E-02	3.59E-01	1 (Unchanged Low
Q9NPJ3 P41226		uncharacterized hypothalamus p		430	443.2575	428.1718	1.09E-01	6.55E-02		Unchanged Low
Q9Y478		ublquitin-activating enzyme E1-lil protein kinase; AMP-activated; b		342	443.334	361.2685	5.70E-01	1.95E-01		Unchanged Low
40.4.0	P06cd8	nodal homolog (mouse)	349,4202	165 252	443.5049 443.7188	304.7576 348.2928	5.35E-01 3.45E-01	-8.93E-01		Unchanged Low
Q12904		small inducible cytokine subfamil		459	445.5171	426.5346	2,50E-01	-4.73E-01 2.94E-01		Unchanged Low
P20264	C02cd1	POU domain; class 3; transcriptic		302	447.8929		-3.44E-02			Unchanged Low
Q15428		splicing factor 3a; subunit 2; 66kl		334	448.6652	355.7289	6.60E-01	2.36E-01		Unchanged Low Unchanged Low
P49771	G06ef6	fms-related tyrosine kinase 3 liga		417	452.1329	439.4747		-1.10E-01		Unchanged Low
P49005	A18cd1	polymerase (DNA directed); delta		232	452,6832	349.3417		-6.49E-01		Unchanged Low
Q99879	M14gh6	H2B histone family; member E	325,9523	483	452.8012	420.7323	4.74E-01	5.69E-01		Unchanged Low
P49821	G20cd1	NADH dehydrogenase (ubiquinos		392	454.6223		-1.70E-01	-3.86E-01		Unchanged Low
P10644	H22ef5	protein kinase; cAMP-dependent	420,8225	518	454.8256	464.5909	1.12E-01	3.00E-01		Unchanged Low
Q9Y5J4	L17gh6	pyrroline 5-carboxylate reductase		303	455,1667			1.06E+00		Unchanged Low
P31040	K02cd2	succinate dehydrogenase comple	671.3369	351	456.2212			-9.36E-01		Unchanged Low
Q9UHA2		synovial sarcoma translocation g		480	456.469	458.4094	5.73E-02	1.30E-01		Unchanged Low
Q9UBF6		ring finger protein 7	315,6541	400	456,9368	390.7196	5,34E-01	3.40E-01		Unchanged Low
Q9Y5S2		CDC42 binding protein kinase be		447	456,9709	430,2759	2.40E-01	2.08E-01		Unchanged Low
P20809	H07ef7	Interleukin 11	399.048	140	458.1544	332.3748	1,99E-01	-1.51E+00	1 (Unchanged Low
Q13526	F21efS	protein (peptidyl-prolyl cis/trans i		460	458.2237	417.3666	4.59E-01	4.66E-01		Unchanged Low
Q9Y6J0	19 H01gh2 E05cd8	hypothetical protein FLJ20671	430.7122	577	458.2293	488.6936	8.93E-02	4.22E-01	11	Unchanged Low
Q9Y4Z6		calcineurin binding protein 1	474.3964	117	458.7489		-4.84E-02			Inchanged Low
Q01433	G02ab2	vacuolar protein sorting 45A (yes	115,16/9	276	458.8586		1,99E+00			Unchanged Low
O431B1		adenosine monophosphate dearr		39D	459.2935	400,6864	3.82E-01			Inchanged Low
	M07ab8 5 D15gh2	NADH dehydrogenase (ubiquinor hypothetical protein FLJ20618	417.4586	432	460.2425	436.6589	1.41E-01	5.03E-02		Jnchanged Low
075431	P19cd6	metaxin 2	354.3599 282.4791	523 443	460.3326	445.8286 395.1258	3.77E-01	5.61E-01		Jnchanged Low
P09543	A18gh7	2;3'-cyclic nucleolide 3' phospho	258 2862	510	460,352 462,033	409.9513	7.05E-01	6.48E-01		Inchanged Low
Q9BTE1		dynactin 4	562,6142	460	462.4054		8.39E-01 -2.83E-01	9.80E-01		Jnchanged Low
P16152	H14ab5	carbonyl reductase 1	234.3184	240	463.6916	312,5841	9.85E-01	3.30E-02		Inchanged Low Inchanged Low
015121	D06cd3	degenerative spermatocyte home		270	464,4612		8.11E-01	3.08E-02		
O60909	E09ab3	UDP-Gai:betaGlcNAc beta 1:4- c		379	464.9697	403.197	3.46E-01	4.98E-02		Jnchanged Low Jnchanged Low
075223	C11gh6	hypothetical protein MGC3077	493,9011	276	465.7709		-8.46E-02			Jinchanged Low
Q9BQA1		MEP50 protein	379.1943	303	466.1997	382.8783	2.98E-01			Inchanged Low
043427	F20cd4	fibroblast growth factor (acidic) in		325	466.8724	371,8858	5.29E-01	6.63E-03		Inchanged Low
	6 A15gh6	hypothetical protein MGC3180	309.3486	596	468.0924	457.7257	5.98E-01	9.45E-01		Inchanged Low
Q9Y6R2		chromosome 4 open reading fran	337.7358	352	469.9502	386,6911	4,77E-01	6.13E-02		Inchanged Low
Q9HBK7		NPD007 protein	367.4893	489	471.1924	442,4205	3.59E-01	4.11E-01		Inchanged Low
P52758	E04ef7	translational Inhibitor protein p14	324.9333	318	471.2741	371.3427		-3.19E-02		Inchanged Low
P47897	E19cd1	glutaminyl-IRNA synthetase	400.1718	364	472.5359		2.40E-01			Inchanged Low
	6 P20gh5	hypothetical protein MGC3262	288.8083			415.0843			1 (Inchanged Low
P35325	H22cd2	small proline-rich protein 2B	390.5307	584		483,0992				Inchanged Low
Q9HA68		hypothetical protein FLJ12154	556,3676	383	474.593		-2.29E-01			Inchanged Low
P49366	P16ab5	deoxyhypusine synthase	458.6886			472.2313				Inchanged Low
Q9UKZ7	J18ef1	colon carcinoma related protein	505.0182	450	478.4787	477.7496	-7.79E-02	-1.67E-01	1 1	Inchanged Low

P01730	K15ef5	CD4 antigen (p55)	516,4046	432	478.8033	475.788	-1.09E-01	-2.57E-01	1	Unchanged Low
	L06cd4	cytochrome c oxidase subunit Va	512.4055	372	480.2708		-9.34E-02			Unchanged Low
Q12828	H02ef6	far upstream element (FUSE) bin	462,592	312	480,5079	418,2868	5.48E-02			Unchanged Low
Q9Y6A4	J18ef3	similar to mouse Git3 or D. malar		485	480,8661	442.0882	4.18E-01	4.32E-01		Unchanged Low
O60888	B09ef2	Other design to the control of the c	298.6981	304	481.136	361.3342 408.0382	6.88E-01	2.62E-02		Unchanged Low
075716	F08cd3	serine/threonine kinase 16	354.1875 327,8121	379 533	481,1652 481,9461	447,6268	4.02E-01 5.56E-01	5.66E-02 7.02E-01	1	
P30711 Q15008	E18ef7	glutathione S-transferase theta 1 KIAA0107 gene product	475,6485	508	482,3327	488.0851	2.01E-02	9.00E-02	1	
P17174	L03gh1 N07ab4	glutamic-oxaloacetic transaminas		224	484.4384	373,5229		-8.84E-01	1	
P13798	K16ab2	N-acylaminoacyl-peptide hydrola		351	484.44	400.3314		-5.51E-02	1	
Q9P2R8	E15ef4	Misshapen/NIK-related kinase	469,5725	511	484.515	488,4865	4.52E-02	1.23E-01	1	Unchanged Low
Q9UNT1	E20cd7	RAB; member of RAS oncogene	268.916	347	484.9919	366.9843	8.51E-01	3.68E-01	1	
O75607	L11cd7	nudeophosmin/nucleoplasmin; 3	350.2387	272	485,5671	369.1972		-3,66E-01		Unchanged Low
O94925	B20gh1	glutaminase	410.4357	471	486.8946	456.0458	2.46E-01	1.98E-01		Unchanged Low
O95390	A06ab3	growth differentiation factor 11	264.606	324 482	489.3912 489.6212	359.3416 498.9664	8.87E-01 -1.01E-01	2.92E-01		Unchanged Low Unchanged Low
O75956	K13cd6	tumor suppressor deleted in oral fally-acid-Coenzyme A ligase; for	524.962 354.1387	174	489.6455	339.2885		-1.02E+00		Unchanged Low
O95573 Q9BXH1	123ab5 N23gh6	Bd-2 blnding component 3	493.3887	433	492.1083	472.8963	-3.75E-03			Unchanged Low
P13662	L20cd5	nudear transport factor 2	392.576	479	492.4942	454.7336	3.27E-01			Unchanged Low
Q92793	H05ab5	CREB binding protein (Rubinstei	468.559	474	494.3525	478.8243	7.73E-02	1.53E-02		Unchanged Low
Q9P0H6	K23ef4	AD-012 protein	406.4789	469	495,1858	456.7944	2.85E-01			Unchanged Low
Q9P0R3	C05ef3	hypothetical protein HSPC213	339.0916	509	495.2656	447.9021	5.47E-01			Unchanged Low
O75094	E11gh1	slit homolog 3 (Drosophila)	345.9855	439	496.3612	426.952	5.21E-01	3.42E-01		1 Unchanged Low 1 Unchanged Low
Q9P011	P10ef2	non-canonical ubquitin conjugati		538 339	497.256 497.6138	498.5084 362.6669	1.11E-01 9.83E-01			Unchanged Low
Q92504 Q95190	M12cd4 L12cd8	HLA class II region expressed ge omithine decarboxylase antizymx		441	497.6839	453,1247	2.41E-01			1 Unchanged Low
Q9Y230	N12cd6	RuvB-like 2 (E. coli)	342,0841	300		379.9805		-1.89E-01		1 Unchanged Low
Q9BSG0	M16gh8	chromosome 2 open reading fran		458		426.9191	6.17E-01	4.98E-01		1 Unchanged Low
Q9Y366	117ef2	chromosome 20 open reading fra		453	498,3057	409,8539	8.41E-01	7.04E-01		1 Unchanged Low
Q14094	O24ef5	cyclin I	407.6242	354	499.6223	420,31		-2.05E-01		1 Unchanged Low
P43403	H04ef5	zeta-chain (TCR) associated pror		396		430.7684		-1.60E-03		1 Unchanged Low
P52803	H23ef7	ephrin-A5	357.5339	316		391.4643		-1.79E-01		1 Unchanged Low
O95402	E18cd5	cofactor required for Sp1 transcri		58,7 399	501.9814 502.2082	222.265 494.287		-8.52E-01 -5.45E-01		1 Unchanged Low 1 Unchanged Low
O00233	L23ab8	proleasome (prosome; macropaia mesoderm development candida:	581.8818 432.735	560		498.6205				1 Unchanged Low
Q9H1K6 Q9BXL6	O18gh7 K15gh6	caspase recruitment domain prot		523	502.8514	483.629	2.42E-01			1 Unchanged Low
Q9H173	H03gh5	endoplasmic reticulum chaperoni		449		469.6418		-2.46E-02		1 Unchanged Low
Q9NV83	H03gh3	hypothetical protein FLJ10876	423.9423	497	504.9702	475.4325	2.52E-01			1 Unchanged Low
P11926	A12ab8	ornithine decarboxylase 1	591,3922	362				-7.07E-01		1 Unchanged Low
Q04743	D23ef1	empty spiracles homolog 2 (Dros		311				-7.73E-01		1 Unchanged Low
Q9BVK2	G11gh6	hypothetical protein MGC2840 si		383				-2.18E-02		1 Unchanged Low
O43251	116ef3	RNA binding motif protein 9	346.6445 355.3783	530 268		461.1452 377.1688	5.49E-01	6.12E-01 -4.09E-01		1 Unchanged Low 1 Unchanged Low
Q99829 Q95777	L13cd4 B10ef2	copine I U6 snRNA-associated Sm-like pr		311		403.8183	3.77E-01			1 Unchanged Low
Q9UN53	C19ef2	calcium binding protein Cab45 pr		407			6.36E-01			1 Unchanged Low
P24310	F03ab4	cytochrome c oxidase subunit VII		288						1 Unchanged Low
Q9H929	N09gh5	hypothetical protein FLJ13055	387.341	500				- 3,67E-01		1 Unchanged Low
P10768	K19ef1	esterase D/formylglutathlone hyd		102				-2,55E+00		1 Unchanged Low
O95287	D17cd5	golgi autoantigen; golgin subfam		560						1 Unchanged Low
Q01105	H18cd1	SET translocation (myeloid leuke		403 551						1 Unchanged Low 1 Unchanged Low
Q9UK45 P35611	B08ef2 117ab2	U6 snRNA-associated Sm-like pr adducin 1 (alpha)	435.0349	519						1 Unchanged Low
O14681	K22cd5	etoposide-induced mRNA	451.5599	324				-4.77E-01		1 Unchanged Low
P80303	A05cd1	nucleobindin 2	296,1418	236				-3.28E-01		1 Unchanged Low
Q9NWC0		F-box only protein 4	434,5394	477	517.7728	476.3476	2.53E-01	1.34E-01		1 Unchanged Low
P00367	B06ab5	glulamate dehydrogenase 1	360.6	303				-2.50E-01		1 Unchanged Low
O00469	109ab8	procollagen-lysine; 2-oxoglulara		238				-1.63E+00		1 Unchanged Low
Q9U104	M18ef4	mitochondrial ribosomal protein i		336				-1.17E-01		1 Unchanged Low 1 Unchanged Low
P20338	N08ab8	RAB4A; member RAS oncogene polymerase (DNA-directed); delta		457 431				-9.49E-02 -2.53E-01		1 Unchanged Low
Q9HCU8 P00519	119gh4 106ef5	v-abl Abelson murine leukemia v		301					•	1 Unchanged Low
Q13880	O02cd5	brain and reproductive organ-exp		263				-1.12E+00		1 Unchanged Low
P25789	L13ef7	proteasome (prosome; macropal		534	526,397	498.9408	2.70E-01			1 Unchanged Low
O43805	J16cd3	Sjogren's syndrome nuclear auto		539						1 Unchanged Low
O75817	A13cd6	POP7 (processing of precursor;		433		445.0255				1 Unchanged Low
Q9UBI1	N11cd7	BUP protein	337,1018	428						1 Unchanged Low
P03950	G12ab2	angiogenin; ribonuclease; RNas		200				1.89E+00		1 Unchanged Low 1 Unchanged Low
P78423 Q9P1D0	F20cd1 E07gh4	small inducible cytokine subfami hypothetical protein PRO2533	58.04051	747 73.3						1 Unchanged Low
P03999	K05ab3	opsin 1 (cone pigments); short-w		37.9				-1.62E+00		1 Unchanged Low
O95793	M22cd2	staufen; RNA binding protein (Di		498				2.54E-01		1 Unchanged Low
P22750	M05ef2	RAB4B; member RAS oncogene		340				2.19E-01		1 Unchanged Low
	B06gh8	ubliquitin UBF-0	431.0731	473	2 540.8482	481.1517		1 1.29E-01		1 Unchanged Low
Q9NRX8	J15gh4	oxidoreductase UCPA	168.7047	52				1.63E+00		1 Unchanged Low
Q9UHQ3		stromal cell protein	469.8784	214				1 -1.13E+00		1 Unchanged Low
Q96S52	J22gh8	phosphatidylinositol glycan; clas MSTP028 protein		28				4.99E-01		1 Unchanged Low 1 Unchanged Low
Q9H3F6 P37198	C24gh8 K03cd8	MSTP028 protein nucleoporin 62kD	442,9007 428,2961	39 49				1 -1.63E-01 1 2.21E-01		1 Unchanged Low
P3/198 Q9H1D4	B05nh4	arginyi aminopeptidase (aminop		49				1 1.47E-01		1 Unchanged Low
075350	K04cd5	glycoprotein; synaptic 2	480.7082	43				1 -1.40E-01		1 Unchanged Low
Q9P024	D16e/3	Huntinglin Interacting protein K	369.5132	48						1 Unchanged Low
Q9Y224	H21ef2	CGI-99 protein	449.089	47			3.05E-0	1 7.30E-02		1 Unchanged Low
Q13361	O12cd3	Microfibril-associated glycoprote		35			1.52E+0			1 Unchanged Low
Q02535	F01ef1	inhibitor of DNA binding 3; domi) 17				0 1.05E+00		1 Unchanged Low
O95864	L24cd4	fatty acid desaturase 2 coatomer protein complex; subu	298.4502	11 47				1 -1.40E+00 1 2.01E-01		1 Unchanged Low 1 Unchanged Low
P53621	H16ab4	coatonici protetti complex; subu	1 409.2/19	41	, 500.031.	, 400.131	· +.34E-0	4.016-01		. One-winged LOW

P35638	N03ab4	DNA-damage-Inducible transcrip	322.5942	318	565.361	401.9484	8.09E-01	-2.12E-02	1	Unchanged	Low
P22307	D22cd1	sterol carrier protein 2	354.3674	356	569.1558	426.421	6.84E-01			Unchanged	
Q92543	D15ght	KIAA0254 gene product	328,3511	356		417.8592	7.95E-01			Unchanged	
P78330	N06ab8	phosphoserine phosphalase	253.4253 401.3697	220	569,9825	472.8205	1.17E+00 5.12E-01		1	Unchanged	
Q9UNX3 P49189	G15ef2 C10ab2	ribosomal protein L26-like 1 aldehyde dehydrogenase 9 famil	446,252	445 277	572,3441 579,314	434.2707		-6.87E-01	1	Unchanged Unchanged	
Canro	H03gh6	inorganic pyrophosphatase	488,3446	402	579,6142	489.9828		-2.81E-01	i	Unchanged	
O15353	H21cd3	winged-hellx nude	407.9375	497	580.6191	495.2368	5.09E-01	2.85E-01	i	Unchanged	
Q9NRP2	D21gh4	DC13 protein	382.8544	421	581.5421	461.731	6.03E-01	1.36E-01	1	Unchanged	Low
Q16280	E18ef1	cyclic nucleotide gated channel a		360	584.8035	469.6748		-3.66E-01	1	Unchanged	
P04183	C04cd2	thymidine kinase 1; soluble	346.7317	289	588.3101	407.8979	7.63E-01		1	Unchanged	
Q12824	O17ef5	SWVSNF related; matrix associa		554	590.2326		1.43E+00		1	Unchanged	
Q99719 P46100	B09ab8 H07ab3	peanut-like 1 (Drosophila) alpha thalassemia/mental relards	419.5269 309.8648	. 410 167	592,2598 594,3041	474,0479 356.9983		-3.19E-02 -8.93E-01	1	Unchanged Unchanged	
Q16610	O05ab5	extracellular matrix protein 1	238,3172	335	596,1691	389.6774	1.32E+00		i	Unchanged	
P39656	E23ab6	dolichyl-diphosphooligosaccharic		367	598.0668	466,4028		-2.39E-01	i	Unchanged	
P53814	G07cd3	smoothelin	517,4102	269	598.16	461.5158		-9.44E-01	1		
Q9P0N4	P24ef1	hypothetical protein LOC51260	609.9142	286	601,9324		-1.90E-02	-1.09E+00	1	Unchanged	Low
Q9BXZ1	A02gh7		465.4301	362	603.8737			-3.62E-01	1	Unchanged	
Q9Y2Q4	H19ef8	HSPC002 protein	337.5615	391	604.9187	444.641		2.14E-01	1	Unchanged	
O15527 Q04323	A18ab8 G01ef2	8-oxoguanine DNA glycosylase ORF	330.4852 497.159	282 255	605.3444	406.0683 454.2076		-2.27E-01 -9.61E-01	1	Unchanged	
P12004	G18ab8	proliferating cell nuclear antigen	332.1465	303	612.821	416.0883		-1.31E-01	1	Unchanged Unchanged	
Q99720	M11cd6	sigma receptor (SR31747 binding		379	615,0729	452,6168	7,57E-01		1	Unchanged	
P43307	G13cd2	signal sequence receptor; alpha	226.1566	322	618.9554	389.0352	1.45E+00	5.10E-01	1	Unchanged	
Q9H3Y8	Q05gh6	chromosome 20 open reading fra		246	619.4944		1.49E+00	1.55E-01	1		
Q9NRX2	A08gh5	mitochondrial ribosomal protein L		305	627,9834				1		
Q9P2X0	L10gh2	dollchyl-phosphale mannosyltran	285.542	431	638,6696		1.16E+00		1	Unchanged	
Q9Y5Z8 P78524	E05cd7 F01cd2	male-specific lethal 3-like 1 (Dro: suppression of tumorigenicity 5	343.4117	394 289	639.7431 643.333	465.0297 425.1425		1.22E-01 -2.50E-01	. 1	Unchanged Unchanged	
P55145	F02ab2	arginine-rich; mutated in early st:		420	656,6324	496.4892	6.68E-01		i	Unchanged	
P33551	O23ab5	CDC28 protein kinase 1	296.3533	355	671,4891	440.861	1.18E+00		i	Unchanged	
Q9BVQ0	E09gh6	hypothetical protein MGC5363	43.30788	54.5	681,2449	259.6855	3.98E+00	3.32E-01	1	Unchanged	
P34897	D15cd2	serine hydroxymethyltransferase	440.9652	290	685.6584	472.1838		-6.05E-01	1	Unchanged	Low
Q9Y5B4	L23ef2	androgen induced protein	268.4081	519	685,9395	491.1814				Unchanged	
Q9Y2V5	B15ef2	transforming growth factor beta 1 solute carrier family 12 (potassiu		327 428	712.3879 729.2688		1.52E+00 1.09E+00		1	Unchanged Unchanged	
Q13953 Q92686	B19cd2 I11ef6	neurogranin (protein kinase C su	330.8792	299	736.2348				1		
Q80565	J05cd8	cysteine knot superfamily 1; BMF	106,6904	151	781.0154	346.3594			i		
Q9Y4M4	121gh3	putative UDP-GaiNAc:polypeptid	346.098	251	846.4309		1.29E+00		1		
P16581	E01ef7	selectin E (endothelial adhesion	109.3094	378	875.1666			1.79E+00	1	Unchanged	
014733	P02ef5	mitogen-activated protein kinase		140	888.014		3.17E+00			Unchanged	
Q16623	M24cd2	syntaxin 1A (brain)	403,2337	165	926.5022			-1.29E+00		Unchanged	
Q9P1M5 O60613	P12gh3 L18cd4	WW domain containing oxidored 15 kDa selenoprotein	38.93 1220.902	19.3 1110	1292.23 1285.07	1205,045		-1.01E+00 -1.38E-01	1	Unchanged Unchanged	
Q16698	M07ab4	2;4-diencyl CoA reductase 1; mit		706		597.2594		5.88E-01	2		
Q9HBA8	D23gh1	24-dehydrocholesterol reductase		278				-1.47E+00	2		
Q9P012	M13ef4	30 kDa protein	546.352	691	614.5795	617.4113			2	Unchanged	Medium
Q9UL53	P24cd7	5'-3' exoribonuclease 2	648.132	604	430.2235			-1.03E-01	2		
P49902	N16cd7	5'-nucleotidase; cytosolic II	1040.523	1220 755		900.9404 697.4428		2.32E-01	2		
O95336 Q03393	P09cd7 A23ab8	6-phosphogluconolactonase 6-pyruvoyltetrahydropterin synth:	524,3104 358,0591	700	812.5955 517.4355	525.0184		5.27E-01 9.66E-01	2	Unchanged Unchanged	
Q13443	P01ef7	a disintegrin and metalloproteina		481	445,3663			-1.44E+00	2		
075173	F15ab2	a disintegrin-like and metalloprot		1060	1254.804			4.29E-01	2		
P51572	O03gh1	accessory proteins BAP31/BAP2	985.1381	889	656,6554	843.6943		-1.48E-01	2	Unchanged	Medium
P42765	D03ab2	acetyl-Coenzyme A acyltransfera		1280	1204.131	1111.329		5.92E-01	2		
P24666	E05ab2 D21cd6	acid phosphalase 1; soluble	1642,968	1970		1833.204		2.59E-01	2		
Q92688 Q14639	B21ab2	acidic (leucine-rich) nuclear phos actin binding LIM protein	1351.973	289 393	835.9103 358.779	531.9498		-7.04E-01 -1.78E+00	2		
O15143	F06ab2	actin related protein 2/3 complex		513				-4.40E-01	2		
O15509	F12ab2	actin related protein 2/3 complex		889	807.751			-3.18E-02	2		
015511	F14ab2	actin related protein 2/3 complex	732.1996	619	466.1372	605.8842	-6.51E-01	-2.42E-01	2		
P12718	A08ab6	actin; gamma 2; smooth muscle;	585.3438	846	2095,96	1175.856		5.32E-01	2		
Q9P016 O43747	O09ef4 M13ab2	AD-015 protein	947.0366	837	1259.571	1014.448		-1.79E-01 -1.05E+00	2		
P20172	A11ab6	adaptor-related protein complex adaptor-related protein complex :	1104.332 1286.83	532 1280				-7.63E-03	2	2 Unchanged 2 Unchanged	
014617	H07ab2	adaptor-related protein complex:		495				-4.57E-01		2 Unchanged	
P07741	M24ab2	adenine phosphoribosyltransfera		862				-1.13E+00	2		
P00568	A02ab2	adenylate kinase 1	2404.119	399		1052.361	-2.76E+00	-2.59E+00		2 Unchanged	Medium
Q9UJJ7	106ef2	adenylale kinase 3 alpha like	758.7288	1060				4.87E-01	2		
P30566	M07ab2	adenylosuccinate lyase	495.8874	622		545.4304		3.27E-01		2 Unchanged	
Q16186	M11cd7	adhesion regulating molecule 1	622.3901 449.3679	610		602.2578 582.9297		-2.95E-02		Unchanged	
Q9NVF6 P26438	B19gh3 B04ab2	ADP-ribosylation factor 1 GTPas ADP-ribosylation factor 6	732.1241	623 818		809.1342		4.72E-01 1.61E-01		 Unchanged Unchanged 	
P40616	D12ab2	ADP-ribosylation factor-like 1	827.1898	1220	1141,92	1064.016		5.64E-01		2 Unchanged	
P36404	D14ab2	ADP-ribosylation factor-like 2	1751.048	1550		1723.74		-1.77E-01		2 Unchanged	
P56559	D20ab2	ADP-ribosylation factor-like 7	485.4562	598	568.6824		2.28E-01	3.00E-01	- 7	2 Unchanged	Medium
P35368	K17ab2	adrenergic; alpha-1B-; receptor	1433,715	1510		1426.069	-1.02E-01	7.34E-02		2 Unchanged	i Medium
P15144	G24ab2 L09ab2	alanyl (membrane) aminopeptida		222				-2.15E+00		2 Unchanged	
P30038 P14550	L05ab2	aldehyde dehydrogenase 4 famil aldo-kelo reductase family 1; mei	1179.797 555.464	598 886		683.9189		-9.81E-01 6.74E-01		2 Unchanged 2 Unchanged	
043488	L07ab2	aldo-keto reductase family 7; mei	546,409	525		606.6855		-5.74E-02		2 Unchanged	
P09972	C16ab2	aldolase C; fructose-bisphosphal	878,5714	1050	947.5062	960.156	1.09E-01	2.63E-01		2 Unchanged	
O43590	G12ef4	alpha-actinin-2-associated LIM p		1070		783.883	-6.31E-01	4.68E-01	- 2	2 Unchanged	Medium
Q08117	M19ab2	amino-terminal enhancer of split	1779.707	1430		1476.318	-5.45E-01	-3.16E-01		2 Unchanged	
Q13438	G05cd7	amplified in osteosarcoma	658.0513	638	802.3407	053.0136	2.86E-01	-4.36E-02		2 Unchanger	Medium

096018	M01gh1	amyloid beta (A4) precursor prob	762.3898	750	1117.582		5.52E-01	-2.38E-02	2 Unchanged Medium
Q13564	P11ab2	amyloid beta precursor protein bi			786.6426		4.21E-01	1.78E-01	2 Unchanged Medium
Q9Y679	B08cd7	ancient ubiquitous protein 1	1191.741	1270	1022.771	1160.442		8.81E-02	2 Unchanged Medium
Q9UKB4	B12ef2	anglomotin like 2	523,7253	776	568,8766		1.19E-01	5.66E-01	2 Unchanged Medium
Q13725	H19ab2	angiotensin receptor 1	1495,332	1360	1058.942	1304.886			2 Unchanged Medium
P08133	120ab2	annexin A6	253.0936	413	879.9474	515.5041		7.08E-01	2 Unchanged Medium
P20073	122ab2	annexin A7	650.1591	481	420.4945	517.0979			2 Unchanged Medium
P03971	E22ab2	anti-Mullerian hormone	1140,812	948	640.7158	909.9247			2 Unchanged Medium
P30041	O08cd5	anti-oxidant protein 2 (non-seleni		1160	3067.945		1.30E+00		2 Unchanged Medium
P27695	K06ef8	APEX nuclease (multifunctional I		653	835.6897		3.27E-01		2 Unchanged Medium
Q9Y4J7	P01ab2	apoptosis inhibitor 5	1850,534	68.4	147.4188	688.7802 -			2 Unchanged Medium
Q9Y5L7	L03ef2	apoptosis related protein APR-3	662.9122	583	1426.489	890.7729			2 Unchanged Medium
O94778	O14ab2	aquaporin 8	1475.036	1420	1474.307		-7.13E-04		2 Unchanged Medium
015296	E02ab2	arachidonale 15-lipoxygenase; s		607	1374.223	718.1334 1288.649			2 Unchanged Medium
P48444	O18ab2	archain 1	1291.871	1200 1600	949,4889		8.92E-02		2 Unchanged Medium
P00966	J24ab2	argininosuccinate synthelase	1049,688	1720	1777.944	1199.548 1518.804	-1.45E-01	6.08E-01 7.10E-01	2 Unchanged Medium 2 Unchanged Medium
P42024	D15ab2	ARP1 actin-related protein 1 hon		724			1.40E-01	4.23E-01	2 Unchanged Medium 2 Unchanged Medium
Q43681	J16ab2	arsA arsenite transporter; ATP-bi		595	509.3934	611.0757			2 Unchanged Medium
Q9UDC3 Q96030	E02ef4 H16ab2	ART-4 protein artemin	728,7274 881,7849	728	638,2016		-4.66E-01		2 Unchanged Medium
P15289	F24ab2	arylsulfalase A	1639.546	1230	1012.121				2 Unchanged Medium
P08243	J18ab2	asparagine synthelase	891.3027	643	1540.113	1024.719	7.89E-01		2 Unchanged Medium
O95630	L17ab2	associated molecule with the SH		872	729.2383		1,43E-01	4.01E-01	2 Unchanged Medium
095135	B09ab2	ataxin 2 related protein	902,4529	345	342,3299	530.0852			2 Unchanged Medium
O14530	N15ab2	ATP binding protein associated v		1390	766.257	993.482	-1.11E-01	7.45E-01	2 Unchanged Medium
P30049	N20ab2	ATP synthase; H+ transporting; r		468	441.4691	511.6658	-5,02E-01	-4.16E-01	2 Unchanged Medium
Q9U112	110ef2	ATPase Inhibitor precursor	776,4612	1220	641.2787	878.5341	-2.76E-01	6.49E-01	2 Unchanged Medium
Q9Y5K8	F17ef2	ATPase: H+ transporting; lysosol	998.0597	1670	1029.886	1232.91	4.53E-02	7.43E-01	2 Unchanged Medium
075787	P15ab2	ATPase; H+ transporting; lysosor		945	1081.688		4.07E-01		2 Unchanged Medium
P05023	P09ef5	ATPase; Na+/K+ transporting; all	1299.236	517	709.0882	841.8422	-8.74E-01	-1.33E+00	2 Unchanged Medium
P45844	P13ef5	ATP-blnding cassette; sub-family	863,8068	381	606.1642	616.8957			2 Unchanged Medium
Q9HAP7	M08gh5	baculoviral IAP repeat-containing		1740	1945,196	1730.54		2.07E-01	2 Unchanged Medium
Q94812	G09ab3	BAI1-associated protein 3	1073.67	983	826.6524				2 Unchanged Medium
P35613	B14ef7	basigin (OK blood group)	603,4821	474	865.2409	647.414		-3.50E-01	2 Unchanged Medium
Q9BSR6	A20gh8	BBP-like protein 1	816,219	1320	914.1927	1015.173	1.64E-01		2 Unchanged Medium
000512	K01ab3	B-cell CLUlymphoma 9	1357,15	76.7	80.36541			-4.15E+00	2 Unchanged Medium
Q12983	E19ab5	BCL2/adenovirus E1B 19kD inter		548	991.4529		1.64E-01	-4.80E-01 4.33E-01	2 Unchanged Medium
Q92934	K24ef1	BCL2-antagonist of cell death	538,8191	727 2510	603.5103 1097.372			2.85E-01	2 Unchanged Medium 2 Unchanged Medium
Q07817 Q43892	E10ef6	BCL2-like 1 Bicaudal D hornolog 1 (Drosophil	2062.673	414	548.7409			-6.63E-01	2 Unchanged Medium
P53004	M23ab3 O19ab3	biliverdin reductase A	490.0724	569	512.2807	523.696		2.15E-01	2 Unchanged Medium
060629	G21ab3	bladder cancer associated protei		1180			8.33E-01		2 Unchanged Medium
Q9HBX3	N21gh5	brain and nasopharyngeal carcin		54				-4.95E+00	2 Unchanged Medium
P56945	H23ef3	breast cancer anti-estrogen resis		1300				-4.14E-02	2 Unchanged Medium
O15255	G21cd6	CAAX box 1	733,6878	577				-3.48E-01	2 Unchanged Medium
P12830	C16ef5	cadherin 1; type 1; E-cadherin (e		924				-2.45E-01	2 Unchanged Medium
P19022	E03ef7	cadherin 2; type 1; N-cadherin (n		540	747.1198	541.7151	1,15E+00	6.79E-01	2 Unchanged Medium
P22223	K19ef7	cadherin 3; type 1; P-cadherin (p		1200	372.3054	823,6929	-1.27E+00	4.22E-01	2 Unchanged Medium
P22676	N07ab5	calbindin 2; (29kD; calretinin)	1781.134	1500				-2,52E-01	2 Unchanged Medium
Q99828	O19gh1	calcium and integrin binding 1 (c	1130.551	1320			-4.31E-01		2 Unchanged Medium
P02593	118ab4	calmodulin 1 (phosphorylase kina		787				-4.51E-01	2 Unchanged Medium
Q13942	L01ab5	calmodulin 3 (phosphorylase kin		464				-9.59E-01	2 Unchanged Medium
P27824	105ef1	calnexin	1492.644	1370				-1.19E-01	2 Unchanged Medium
P17655	C20ef6	calpain 2; (m/ll) large subunit	671.5939	837			-2.90E-01		2 Unchanged Medium
P04632	C22ef6	calpain; small subunit 1	1565.935	1650 603			-3.16E-01		2 Unchanged Medium
Q15417	P17ab5	calponin 3; acidic	911.495 629.0213	486				-5.95E-01 -3.73E-01	2 Unchanged Medium 2 Unchanged Medium
Q9UEG5 P52907	D17ef8 O08ab4	capicua homolog (Drosophila) capping protein (actin filament) n		684				-3.11E-01	2 Unchanged Medium
Q9GZX3	N16gh1	carbohydrate (N-acetylglucosam		929			-7.81E-02		2 Unchanged Medium
Q16619	K01ab4	cardiotrophin 1	1478.249	1220				-2.72E-01	2 Unchanged Medium
Q9UBD9	124cd8	cardiotrophin-like cytokine; neuro		893				1,30E+00	2 Unchanged Medium
Q15699	M16cd4	cartilage paired-class homeoprol		596				-3.26E-01	2 Unchanged Medium
P48729	A22ab6	casein kinase 1; alpha 1	690,8551	1150	902.9474	914,9218	3.86E-01	7.36E-01	2 Unchanged Medium
P48730	L07ab4	caseln kinase 1; delta	951.5238	1160	1111.571	1073,796	2.24E-01	2.84E-01	2 Unchanged Medium
P19138	L08ef5	casein kinase 2; alpha 1 polypep	483,6062	597	461.8036	513,9702	-6,66E-02	3.03E-01	2 Unchanged Medium
P49662	C12ef6	caspase 4; apoptosis-related cys	2013.957	2460	1017.425	1831.485	-9.85E-01	2.90E-01	2 Unchanged Medlum
P21964	F19ef6	catechol-O-methyltransferase	697.5886	1170					2 Unchanged Medium
P26232	K01ef7	catenin (cadherin-associated pro		717	810.257			-1.54E-01	2 Unchanged Medium
P35222	G07ef7	catenin (cadherin-associated pro		1290					2 Unchanged Medium
P07339	N23ef7	cathepsin D (lysosomal asparty)		947				-4.04E-01	2 Unchanged Medium
Q99967	C18cd6	Cbp/p300-Interacting transactiva		1600					2 Unchanged Medium
P49715	N17ef6	CCAAT/enhancer binding protein		884		937,3612			2 Unchanged Medium
P53567	N22ab5	CCAAT/enhancer binding protein		1160				4.92E-01	2 Unchanged Medium 2 Unchanged Medium
095627	B21cd8 G02cd4	CCR4-NOT transcription compte		851 570				4.89E-01 -3.97E-01	2 Unchanged Medium
Q04900	P07ab3	CD164 antigen; sialomucin CD3E antigen; epsilon polypepti	750,3086	1410				-3.59E-01	2 Unchanged Medium
P07766 P34810	A21ab4	CDSE antigen; epsilon polypepii CD68 antigen	2460.227	1100				-1.16E+00	2 Unchanged Medium
P18582	B21ab5	CD81 antigen (target of antiproli		938				-4.75E-01	2 Unchanged Medium
Q16181	F09ef5	CDC10 cell division cycle 10 hor		443				5.42E-02	2 Unchanged Medium
Q16543	H01ef5	CDC37 cell division cycle 37 hor		1050				-4.35E-02	2 Unchanged Medium
014735	114cd6	CDP-diacylglycerol-inositol 3-pl		1180		579,4931		2.29E+00	2 Unchanged Medium
P30260	F13ef5	cell division cycle 27	1060,812	1470				4.74E-01	2 Unchanged Medium
P49427	B12ef7	cell division cycle 34	762.5808			942.0784		7.66E-01	2 Unchanged Medium
P29373	F01ef7	cellular retinoic acid binding pro		803				-6.11E-01	2 Unchanged Medium
P41208	F09ab5	centrin; EF-hand protein; 2	539,1633	672		2 629,3345	3,28E-01	3.18E-01	2 Unchanged Medium
000522	F05ab4	cerebral cavernous malformation	n 870.4309	631	723.1482	2 741.4854	-2.67E-0	4.64E-01	2 Unchanged Medium

015183			402,949	610	0 883,328	13 632 0801	8 1.13E+00		- 1
Q9Y2Z		CGI-06 protein	721.6709	67				0 5.98E-01 1 -9.94E-02	- orientaliged inscalati
Q9Y3C		CGI-120 protein	1229,537	136	0 2124.30	4 1572.302			2 Unchanged Mediun 2 Unchanged Mediun
Q9Y3E		CGI-128 protein CGI-147 protein	1592.767	2370			2 3.37E-01	5.75E-01	2 Unchanged Medium
Q9Y3E			532.425 785.276		8 567,063		7 9.09E-02	2 5.66E-01	2 Unchanged Medium
Q9Y320			745.0314	924 933		9 829.1206	-1.33E-02	2.35E-01	2 Unchanged Medium
Q9Y357		CGI-40 protein	817.7272	769			8.22E-02	3.22E-01	2 Unchanged Medium
Q9Y360		CGI-45 protein	786,0567	914				-8.87E-02 2.17E-01	2 Unchanged Medium
Q9Y512		CGI-51 protein	967.3508	859				-1.71E-01	2 Unchanged Medium
Q9Y390 P40227			1391,902	802			2 -2.88E-01	-7.96E-01	2 Unchanged Medium 2 Unchanged Medium
Q99832	H07cd6		1011.613	436		4 679.9043	3 -7.72E-01	-1.22E+00	2 Unchanged Medium
Q9UBR		chaperonin containing TCP1; sul chemokine-like factor 1		1090		4 1274.983	3.79E-01	-1.33E-01	2 Unchanged Medium
Q9UHN		chemokine-like factor 1	1047.63	1800				7.85E-01	2 Unchanged Medium
P54105		chloride channel; nucleolide-sen	1568,851	2030 487			-3.42E-01		2 Unchanged Medium
Q14781		chromobox homolog 2 (Pc dass)	1 795.873	749					2 Unchanged Medium
Q9BXS4		chromosome 1 open reading fram	874.0189	1370				-8.68E-02 6.52E-01	2 Unchanged Medium
Q9NPAC		chromosome 11 hypothetical pro	998.8577	834			-2.18E-01	-2 605-01	2 Unchanged Medium
Q9UKR5 P56378			1181.962	1160				-3.18E-02	2 Unchanged Medium 2 Unchanged Medium
O95433			503.5233	690					2 Unchanged Medium
Q9NV31		chromosome 14 open reading fra chromosome 15 open reading fra	656.1775	563			-5.80E-01	-2.20E-01	2 Unchanged Medium
Q9NVE2		chromosome 2 open reading fran	648.49	832				·3.76E-02	2 Unchanged Medium
Q9U105	L04gh4	chromosome 20 open reading fra	2518 951	671 2140		2 596.0999	-4.69E-01	4.96E-02	2 Unchanged Medium
Q9P0A7		chromosome 20 open reading fra	581,9098	1080			-1.07E+00	-2.36E-01	2 Unchanged Medium
Q9Y3B1		chromosome 20 open reading fra	1691.864	1230		1235 108	1.04E+00 -1.11E+00	8.93E-01	2 Unchanged Medium
Q9BQ89		chromosome 20 open reading fra	1460.875	952			-2.58E-01	-9.00E-01	2 Unchanged Medium
Q96C58 Q9Y5Z4		chromosome 6 open reading fran	1254.647	1080		940.0036	-1.38E+00	-2 10F-01	2 Unchanged Medium
Q00610	K12ef3 H2cd5	chromosome 6 open reading fran		988		758.6366	-1.30E+00	1.10E-01	2 Unchanged Medium 2 Unchanged Medium
P56749	F04cd7	clathrin; heavy polypeptide (Hc) claudin 12	1639,759	1210	1437.44	1427.503	-1,90E-01	-4.44E-01	2 Unchanged Medium
O43809	M15cd7	cleavage and polyadenylation sp	2262,278	2480		1992.755	-8.75E-01		2 Unchanged Medium
O96005	G05ab4	cleft lip and palate associated tra	1353 338	503 1170					2 Unchanged Medium
Q16740	E20cd4	ClpP caseinolylic protease; ATP-	625.8711	528	475.0848		4.36E-02 -3.98E-01	-2.15E-01	2 Unchanged Medium
P55085	O24ab5	coagulation factor II (thrombin) re	829.7537	979	392.7495		-1.08E+00	2.39E-01	2 Unchanged Medium
P13726 Q9Y678	805ab4 D14ef8	coagulation factor III (Ihrombopla		880		800.9723	-1.29E+00	-2.94F-01	2 Unchanged Medium 2 Unchanged Medium
Q15363	M11ef3	coat protein gamma-cop	815.7993	738	1009,756	854,5161	3.08E-01	-1.45E-01	2 Unchanged Medium
043513	N06cd4	coated vesicle membrane proteir cofactor required for Sp1 transori	706.3054 504.8040	804	870.2118		3.01E-01	1.88E-01	2 Unchanged Medium
Q9Y281	N18cd8	cofilin 2 (muscle)	671.9394	481	798.8706		4.33E-01		2 Unchanged Medium
P04141	G22gh6	colony stimulating factor 2 (grant	116.2139	270 1290	1093.534 165.718		7.03E-01		2 Unchanged Medium
P02745	L06ef1	complement component 1; q sub-	683,4431	1650	798.2366	522.783 1042.465	5.12E-01		2 Unchanged Medium
P29279	116ab4	connective tissue growth factor	596,4123	537	1071.517		2.24E-01 8.45E-01		2 Unchanged Medium
O43191 Q99627	B02cd3	COP9 constitutive photomorphos	855.8616	1030	1302,334			2.68E-01	2 Unchanged Medium 2 Unchanged Medium
Q35027 Q15387	C01cd7 G21cd7	COP9 homolog	788.2297	968	1195.497	984.0834	6.01E-01		2 Unchanged Medium 2 Unchanged Medium
Q9ULV4	K18ef3	COP9 subunit 6 (MOV34 homolo coronin; actin binding protein; 10	590.9973	382	628.5041		8.88E-02	6.30E-01	2 Unchanged Medium
Q98R76	114gh7	coronin; actin-binding protein; 18	708 5657	1200 1080	1362.545	1385.839	-2.26E-01	-4.09E-01	2 Unchanged Medium
Q14061	B22cd5	COX17 homolog; cytochrome c c	889.1581		690.8568 745.8658	827.7864	-3.65E-02	6.13E-01	2 Unchanged Medium
Q9Y6B2	E02ef8	CREBBP/EP300 inhibitory protel	946.9513	1030	1988.725	1026,374 1323,308	-2.54E-01	7.00E-01	2 Unchanged Medium
Q9NY68	C06gh4	CTL2 gene	1381 821			991.4995	-R 09F-01 .	7.2/E-01	2 Unchanged Medium
P24385 Q9NXT4	004ef5 K09gh2	cyclin D1 (PRAD1: paralhyroid a		2150	1100.478	1486.769	-1.42E-01	8.21E-01	2 Unchanged Medium 2 Unchanged Medium
P01034	111ab6	cyclin M2	1560.659		1582.768	1620.99	2.03E-02	1.40E-01	2 Unchanged Medium 2 Unchanged Medium
Q15828	113ab4		1067,127		1366.396	1134.612	3.57E-01 -	1.37E-01	2 Unchanged Medium
P01036	C02ab6	andali - A	967,1669 1261,605		451,3932	1404.267 -	1.10E+00	1.53E+00	2 Unchanged Medium
P52943	105ab4	cysteine-rich protein 2	B74.1939		1069.756 1277.819	932.4892 -	-2.38E-01 -	1.44E+00	2 Unchanged Medium
P13498	G18ab3	cytochrome b-245; alpha polyper	708.9954			883.82 707.7483 -	5.48E-01 -	8.08E-01	2 Unchanged Medium
P00167 P00001	C04ab6	cytochrome b-5	2187.128		1034,311	1565.893 -	3.81E+01	0./3E-01 5.67E-01	2 Unchanged Medium
P10606	N24gh6 124gh6	cytochrome c	547.6708			502.4786 -	8.14E-02	3.09E-01	2 Unchanged Medium 2 Unchanged Medium
P08574	P10ab5	cytochrome c oxidase subunit Vb cytochrome c-1	1364.01		2087.456	1851.225	6.14E-01	6.24E-01	2 Unchanged Medium
Q99426	C17gh1	cytoskeleton-associated protein	567.9346	443	441.6771	517.4383 -	5.97E-01 -	5.83F-01	2 Unchanged Medium
Q07065	G13cd7	cytoskeleton-associated protein · 6	500.9411	684 575	177.3269	673.9833	4.71E-01	2.85E-01	2 Unchanged Medium
Q1503B	L13ef3		308.0755		564 7611	987.5887 1	1.57E+00 -		2 Unchanged Medium
Q9H2L4	B18gh8	DC32	08.0427	702	780.2878	754.4305 -	5.1/E-01 ·	1.40E-01	2 Unchanged Medium
Q9NPA8	G22gh7	DC6 protein c	MONE MAG			630.0414 ± 1342,189 ±		7.82E-01 7.32E-01	2 Unchanged Medium
060231 Q9GZR7	H13cd3	DEAD/H (Asp-Glu-Ala-Asp/His) t 7	53.8815	873	799.6561	808,7915	R 50F.02 1	145.04	2 Unchanged Medium 2 Unchanged Medium
P17844	E02gh4 F04ab4	DEAD/H (Asp-Glu-Ala-Asp/His) t	803.42	984	774.3765	854.0965 -	5315-02 3	035.04	2 Unchanged Medium 2 Unchanged Medium
Q08211	M03ab4	DEAD/H (Asp-Glu-Ala-Asp/His) t 1	482.642	1440	10/0.104	153D 749 4	1 72E_n+ ./	200.02	2 Unchanged Medium
G9UJV9	D08ef2	DEAD/H (Asp-Glu-Ala-Asp/His) t 9 DEAD-box protein abstrakt		628 8	907,4343	B24.2193 -2	2.41E-04 -4	.65E-01	2 Unchanged Medium
P51398	G18ef6	death associated protein 3	76.8378 20.3044	1230 (0/0.36/2	1024.437 -1	1.66E-01 3	1.28E-01	2 Unchanged Medium
Q13437	G22cd4	Deleted in split-hand/split-foot 1 i 1	565.344	1500 1	1721 P72	905.1904 -1 1596.672 1	1.76E-01 9	.20E-02	2 Unchanged Medium
060735	O06cd6	dendritic cell protein 8	58.6375	502 4	167.3432	609,2302	1.3/E-01 -5 3.78E 04 -	.68E-02	2 Unchanged Medium
Q16854	M06ab5	deoxyguanosine kinase a	58 1918			609.2393 -6 910.8082 2	2.05F-01 4	.13E-01	2 Unchanged Medium
Q9Y295 P00387	H04ab8 F09ab3	developmentally regulated GTP 1 7	36.3786	634 E	000.7023	576.9267 - 1	1,56E-01 -2	.17F-01	2 Unchanged Medium
	108ef2	diaphorase (NADH) (cytochrome 1	402.134	1030	1435.74	1290.593 3	3.42E-02 -4	.40E-01	2 Unchanged Medium 2 Unchanged Medium
	C06ab6	dicarbonyl/L-xylulose reductase 5	44.2125	740 9	538.7BU7	/4117	7 Q7 C N4 4	.44E-01	2 Unchanged Medium
	M12ef7	dihydrolipoarnide S-succinyltrans 6: diphtheria toxin receptor (heparir 7:	23,503/ 21.8424	/13 3	93.7165 (577.6196 -6	69F_01 1	BBC.A+	2 Unchanged Medium
Q9UFN1	H10gh6	DKFZP564B147 protein 9	21.8431 66.5927	638 947 1	187.117	518.9525 -1.	.87E+00 -1	.76E-01	2 Unchanged Medium
	C13ef8	DKFZP564C1940 protein 9	41.4076	865	804.639	1127.658 6	.04E-01 -2	.91E-02	2 Unchanged Medium
	C07ef8	DKFZP564M082 protein 6	75.8991			970.3185 -2 998.4411 B	20E-01 -1	.2ZE-01	2 Unchanged Medium
Q9NWE3	Jilef8		19.5607			605.084 5	1.38E-01 7	.1/E-U] 76E-04	2 Unchanged Medium
								.r 0E-U }	2 Unchanged Medium

Q9H6Y7	H04gh6	DKFZP566H073 protein	1390,389	1480	1191.195	1352.292	-2.23E-01	8.55E-02	2 Unchanged Medium
Q9Y3X0	P23ef8	DKFZP586M1019 protein	588.3603	505	672.4872	588.5767	1.93E-01	-2.21E-01	2 Unchanged Medium
060762	G19cd4	dollchyl-phosphate mannosyltran	407.0059	581	605.8353	531.3373	5.74E-01	5.14E-01	2 Unchanged Medium
P53805	B11ab5		370.0907	663	511.9248	515.0794	4.68E-01	8.42E-01	2 Unchanged Medium
Q13597	N13el3	downregulated in ovarian cancer	594,6931	693	623.702	636,966	6.87E-02	2.20E-01	2 Unchanged Medium
Q9C005	J05gh8	dpy-30-like protein	810.4585	1210	1174.001	1064,132	5.35E-01	5.76E-01	2 Unchanged Medium
Q13448	J03cd6	DR1-associated protein 1 (negati	1240.359	1140	2382.895	1586.247	9.42E-01		2 Unchanged Medium
Q9UHF9	K01ef4	dual oxidase 2	1146.14	648	316.6462		-1.86E+00		2 Unchanged Medium
P51452	N22ab4	dual specificity phosphatase 3 (v		605	473.5652		-4.56E-01		2 Unchanged Medium
Q14203	A13ab6	dynactin 1 (p150; glued homolog		529	623.6948		2.97E-01	6.06E-02	2 Unchanged Medium
Q13561	D19cd6	dynactin 2 (p50)	1565.567	1380	1173.024	1372.261		-1.84E-01	2 Unchanged Medium
Q13409	116ef1	dynein; cytoplasmic; intermediate		604	464.9675		-7.71E-01		2 Unchanged Medium
Q09472	C18ab4	E1A binding protein p300	1496.409	2120	1342.944	1651.644		5.00E-01	2 Unchanged Medium
	D20gh8	EAF1 protein	642.1587	652	469.6834	588.0816		2.28E-02	2 Unchanged Medium
P78365	N19ab4	early development regulator 2 (p		1070	761.5923	907.5945	-2.56E-01	2.54E-01	2 Unchanged Medium 2 Unchanged Medium
Q9984B	G11cd7	EBNA1 binding protein 2	977.8671	611 332	818.6583 458.9563		-1.13E+00		2 Unchanged Medium 2 Unchanged Medium
Q12805	A21ab6	EGF-containing fibulin-like extrat	1002.82	500	629,1845		-2.28E-01		2 Unchanged Medium
Q9Y6I3	D21cd8 C04ab4	EH domain-binding mitotic phosp ELAV (embryonic lethal; abnorms		800	843.4463		-1.26E-02		2 Unchanged Medium
Q15717 P13804	124ab3	electron-transfer-flavoprotein; alt		545	529.2866		-7.96E-01		2 Unchanged Medium
Q14247	K17ef6	ems1 sequence (mammary lumo		1050	1002.084		-1.80E-01		2 Unchanged Medium
P42892	O21ab4	endolhelin converling enzyme 1	602.4294	799	809,7071	737.1982	4.27E-01	4.08E-01	2 Unchanged Medium
P30084	A17ab6	encyl Coenzyme A hydratase; sh		1210	741,2226		-1.14E-01	5.93E-01	2 Unchanged Medium
P07099	120ab3	epoxide hydrolase 1; microsomal	1231.48	1840	1198.169	1424.661	-3.96E-02	5.83E-01	2 Unchanged Medium
P29692	H04ab5	eukaryotic translation elongation	1329.756	744	1110.87	1061.631	-2.59E-01	-8.37E-01	2 Unchanged Medium
P20042	L05cd4	eukaryotic translation initiation fa		603	956,276	1016.24	-6.39E-01	-1.30E+00	2 Unchanged Medium
P41091	A20ab4	eukaryotic translation initiation fa	742,5843	474	478.7249	565.2535			2 Unchanged Medium
Q99613	L16cd3	eukaryotic translation initiation fa		853	998,7488		-1.91E-01		2 Unchanged Medium
P78344	C02ab4	eukaryotic translation initiation fa	1562.952	1520	1177.845			-4.19E-02	2 Unchanged Medium
P23588	A24ab4	eukaryotic translation initiation fa		1010	1125.825		-2.54E-01		2 Unchanged Medium
Q13541	C03ab5	eukaryotic translation initiation fa		1060	2293,236	1669.873		-6.43E-01	2 Unchanged Medium
Q13542	A19ab6	eukaryotic translation initiation fa		1850	1663,197	1712.605	3.79E-02		2 Unchanged Medium
O60573	G16cd5	eukaryotic translation initiation fa		883	819.3679	833,9147	3.49E-02		2 Unchanged Medium
Q16394	K02ab3	exostoses (multiple) 1	513,7526	428	813.2693	584,9471		-2.64E-01	2 Unchanged Medium
P37268	C17ab6	farnesyl-diphosphate farnesyltrat		455 771	649,0518 476,1039	610.2023 566.0193	7.86E-02	-6.73E-01 7.74E-01	2 Unchanged Medium 2 Unchanged Medium
P55899	L22ab5	Fc fragment of IgG; receptor; tran	548.2677	547	530.801	541,9489		-3.93E-03	2 Unchanged Medium 2 Unchanged Medium
Q9UK73	P05ef3 G14ab4	fem-1 homolog b (C. elegans) filamin A; alpha (actin binding pro		675	1277.458	966,2982		-4.86E-01	2 Unchanged Medium
P21333 P26885	P07ab4	FK506 binding prolein 2 (13kD)	814.6357	510	661,4593		-3.01E-01		2 Unchanged Medium
Q13451	F15ab4	FK506 binding protein 5	1366.905	1110				-2.98E-01	2 Unchanged Medium
095633	M01cd6	follistatin-like 3 (secreted glycopr		1280	1092.99	1227.557	-2.64E-01		2 Unchanged Medium
P15407	A20cd4	FOS-like antigen 1	1696.135	1350	1218.491	1421.01			2 Unchanged Medium
Q14192	G02ab4	four and a half LIM domains 2	309.9219	591	1210.635	703.7424	1.97E+00	9.30E-01	2 Unchanged Medium
Q9NZA0	N17gh6	fuse-binding protein-interacting r	924.2347	911	1110,468	981.791	2.65E-01	-2.13E-02	2 Unchanged Medium
P35637	P19ab4	fusion; derived from t(12;16) mall	866.3996	654	1159,792		4.21E-01	-4.05E-01	2 Unchanged Medium
	E21gh7	FXYD domain-containing ion tran		1060			-2.32E-01	1.71E-01	2 Unchanged Medium
Q9Y653	B02cd5	G protein-coupled receptor 56	2010,489	2160		1889.081		1.00E-01	2 Unchanged Medium
Q9H1C0	E18gh4	G protein-coupled receptor 92	712.1007	750		727.2869	1.55E-02		2 Unchanged Medium
P15170	L13ab4	G1 to S phase transition 1	2042.425	1410		1521,226			2 Unchanged Medium
095166	D09cd7	GABA(A) receptor-associated pro		2210		1572.36			2 Unchanged Medium
008765	D21cd7	GABA(A) receptor-associated pri		1130 800		1095.839 665.3025		-1.39E-01 5.69E-01	2 Unchanged Medium 2 Unchanged Medium
P16278	O05ef1	galactosidase; beta 1 GCN5 general control of amino-a	539.4222	637	919,5196			-3.52E-01	2 Unchanged Medium
P78537	J23ab4 F14ab4	GDP dissociation inhibitor 1	633.6157	894	640.108				2 Unchanged Medium
P31150 P06396	B23ab3	gelsolin (amyloidosis; Finnish typ		577					2 Unchanged Medium
P52657	E03ab6	general transcription factor IIA; 2		685					2 Unchanged Medium
Q00403	F08ef6	general transcription factor IIB	500.4565	791	454.8328	582,2035	-1.38E-01	6.61E-01	2 Unchanged Medium
P48060	123cd7	glioma pathogenesis-related pro		692					2 Unchanged Medium
P06744	B19ab3	glucose phosphale isomerase	900.9772	553	726,4749	726,7688	-3.11E-01	-7.05E-01	2 Unchanged Medium
P30101	G13ab5	glucose regulated protein; 58kD	1708.658	1250	2220.105			-4.50E-01	2 Unchanged Medium
043836	122ef1	glutamate receptor; ionotropic; N		1890			-6.36E-02		2 Unchanged Medium
P15104	H22ab5	glutamate-ammonia ligase (gluta		1040				-1.71E-01	2 Unchanged Medium
Q06210	A23ab5	glutamine-fructose-6-phosphate		600				-5.34E-01	2 Unchanged Medium
Q9Y3D4		glutaredoxin 2	888.5406	894				8.32E-03	2 Unchanged Medium
Q9Y2Q3		glutathione S-transferase subun		346				-1.28E+00	2 Unchanged Medium
P48637	D01ab3	glutathione synthetase	650.8496	429			-4.13E-01	-6.00E-01 2.95E-02	2 Unchanged Medium
P78417	E20cd5 J14ef4	glutathione-S-transferase like; gl		2020 1460					2 Unchanged Medium 2 Unchanged Medium
P49840		glycogen synthase kinase 3 alph	1 1037,300 - 407,4684	530					2 Unchanged Medium
P13224 Q9UBQ7	F23ab3 L08cd7	glycoprotein ib (platelet); beta po glycxylate reductase/hydroxypyr	947.0062	977				4.55E-02	2 Unchanged Medium
Q9P2T1	C02ef2	GMPR2 for guanosine monopho		1070					2 Unchanged Medium
Q9H4A6		golgi phosphoprotein 3 (coat-pro		1210				2.35E-01	2 Unchanged Medium
P28799	B19ab5	granulin	1611.298	1750			-3.79E-01		2 Unchanged Medium
Q14393	H10ab3	growth arrest-specific 6	484.489	563					2 Unchanged Medium
P29354	H06ef5	growth factor receptor-bound pre		854			-1.22E-01		2 Unchanged Medium
Q14451	D20ab4	growth factor receptor-bound pro	712.3296	846		731.6029	-1.61E-01	2.47E-01	2 Unchanged Medium
Q9Y6G2		growth hormone inducible transp	1289.256	1020				-3.43E-01	2 Unchanged Medium
P04899	J09ab4	guanine nucleolide binding prote	850.9208	1390				6.94E-01	2 Unchanged Medium
P11016	M13ef5	guanine nucleotide binding prote	e 758.629	773					2 Unchanged Medium
Q13905	G09ab6	guanine nucleotide-releasing fac		569					2 Unchanged Medium
Q16774	L16ab3	guanylate kinase 1	968,5867	1340				4.72E-01	2 Unchanged Medium
Q9Y649	K16el4	GW128 protein	1663.305	2420					2 Unchanged Medium
P07305	G13ab6	H1 histone family; member 0	1780.353	1870				6.73E-02	2 Unchanged Medium 2 Unchanged Medium
075367	M14cd5	H2A histone family; member Y	1065.311	837 461				-3.48E-01 -3.58E-01	2 Unchanged Medium
P17317	C24ab6	H2A histone family; member Z	591.0013 586.8773	85				5.38E-01	2 Unchanged Medium
P02278	M16gh6	H28 histone family; member G	200.0113	0.5		, 01,31	ZZE-U		- summinged mediting

	Q9NS37	A03gh5	HCF-binding transcription factor.	857,3143	717	601.7268	725.1835	-5.11E-01	-2 59F-01	2 Unchanged Medium
	O96004	E02cd5	heart and neural crest derivatives		612	621.033		-1.84E-01		2 Unchanged Medium
	P10809	M11ef5	heat shock 60kD protein 1 (chap-	1693.23	1020	1276.41		-4.08E-01		2 Unchanged Medium
	P34932	111ef1	heat shock 70kD protein 4	1402,505	1610	1042.113		-4.28E-01		2 Unchanged Medium
	P07900	G18ef7	heat shock 90kD protein 1; alpha	2405.793	1430	1512.258		-6.70E-01		2 Unchanged Medium
	P08238	B20ab7	heal shock 90kD protein 1; beta	2781.462	1950	979.1145		-1.51E+00		2 Unchanged Medium
	O75506	H17ab6	heat shock factor binding protein	1132,126	1300	1081,748		-6.57E-02	1,99E-01	2 Unchanged Medium
	Q9UHG4	K18ef4	heme-regulated initiation factor 2	413.2841	659	576.3971	549,5035	4.80E-01	6.73E-01	2 Unchanged Medium
	P09105	J21ab4	hemoglobin; theta 1	702.445	484	769,2336	651.9692		-5.37E-01	2 Unchanged Medium
	O43504	D23cd6	hepalitis 8 virus x Interacting pro		1340	949.0116	.1097.672	-8.79E-02		2 Unchanged Medium
	Q9H2I6	N10gh5	hepailtis C virus core-binding pro	676.7297	620	473.9328		-5.14E-01		2 Unchanged Medium
	P51858	K16ef7	hepatoma-derived growth factor i		1170	490.7456	850.7213			2 Unchanged Medium
	Q04150	A08ab7	heterogeneous nuclear ribonucle		870	1065,597	959,787	1.76E-01	-1.16E-01	2 Unchanged Medium
	Q13151	E15cd7	heterogeneous nuclear ribonucle	906.986	944	1143.552	998,1703	3.34E-01	5,77E-02	2 Unchanged Medium
	P22626	N09ab6	heterogeneous nuclear ribonucle	1021.87	1010	1584.855	1206,298	6.336-01	-1.38E-02	2 Unchanged Medium
	P07910	A10ab7	helerogeneous nuclear ribonude		718	1059.349	841.6601		-5.79E-02	2 Unchanged Medium
	O14979	L09cd5	heterogeneous nuclear ribonucle		1990	1646.172	1563,717	6.38E-01	9.10E-01	2 Unchanged Medium
	P52597	G06ab7	heterogeneous nuclear ribonucle		1280	1517.274	1567.536		-5.69E-01	2 Unchanged Medium
	Q07244	N13ab6	heterogeneous nuclear ribonucle		667	684.4978	733.1055			2 Unchanged Medium
	Q15584	P03ab7	helerogeneous nuclear ribonucle		569	858.2829	717.4409		-3.51E-01	2 Unchanged Medium
	P12081	J12ab4	histidyl-tRNA synthetase	555.5578	565	424.3904	514.8575			2 Unchanged Medium
	Q9UBN7 Q13838		histone deacetylase 6	685.5336	407	695,1552	595.8457		-7.53E-01	2 Unchanged Medium
		M07cd4	HLA-B associated transcript 1	681.4666	604	921.7532	735.5784		-1.75E-01	2 Unchanged Medium
	Q99873 Q9H063	H02ef7	HMT1 hnRNP methyltransferase	890.1184	836	1209.872	978.6584		-9.05E-02	2 Unchanged Medium
	P51610	M14gh8	homolog of yeast MAF1	744.9951	921	1013.86	893,139		3.05E-01	2 Unchanged Medium
	O00165	L21ab5 I04cd6	host cell factor C1 (VP16-access		580	735.1691	632,9786		-9.25E-03	2 Unchanged Medium
	Q9Y2Q5	P24ef7	HS1 binding protein	1482.068	1490	2425.818	1800.489	7.11E-01	1.12E-02	2 Unchanged Medium
	Q9Y2R0	A05ef8	HSPC003 protein	620,7783	1010	625.6574	752,165	1.13E-02		2 Unchanged Medium
	Q9Y2TO	A17el8	HSPC009 protein HSPC022 protein	1507.005	1720	1784.568	1668.909	2.44E-01	1.87E-01	2 Unchanged Medium
	Q9UNZ5	C17ef8	HSPC023 protein	453.0964 825.3888	733	326,4215		-4.73E-01	6.94E-01	2 Unchanged Medium
	Q9P032	019ef8	HSPC125 protein	558,8905	1310	1347.488	1159.538	7.07E-01	6.62E-01	2 Unchanged Medium
	Q9P019	A06ef8	HSPC141 protein	717.9096	1700	843,0953 1519,292	668.1435	5.93E-01	1.08E-01	2 Unchanged Medium
	Q9P004	A24ef8	HSPC160 prolein	1182.006	1070	545.8523		1.08E+00 -1.11E+00		2 Unchanged Medium
	Q9P003	C04ef8	HSPC163 protein	801.5204	1410	1148.378	1118,827	5.19E-01	8,11E-01	2 Unchanged Medium
	Q9P000	C06ef8	HSPC166 protein	508.2074	921	523.3806	650.7428	4.24E-02	8.57E-01	2 Unchanged Medium
	Q9Y684	N09ef2	HSPCO34 protein	569,9533	628	797.4819	665.044	4.85E-01	1.39E-01	2 Unchanged Medium 2 Unchanged Medium
	Q9NRG2	D01gh4	HTGN29 protein	1276.132	1470	1639,451	1462.8	3.61E-01	2.07E-01	2 Unchanged Medium 2 Unchanged Medium
	Q16775	G15ab6		511.3412	676	407,0538		-3.29E-01	4.03E-01	
	P40939	D07ab3		1814.498	1770	1617.578		-1.66E-01		2 Unchanged Medium 2 Unchanged Medium
	P55084	D09ab3	hydroxyacyl-Coenzyme A dehydr	539,2691	890	1218,105		1.18E+00		2 Unchanged Medium
1	Q14526	B13ab5	hypermethylated in cancer 1	918,3678	904	1008.759		1.35E-01		2 Unchanged Medium
	Q9UBS2	A15ef1	hyperpolarization activated cyclic	676,9289	688	520.6835		-3.79E-01		2 Unchanged Medium
	Q96EW8	P08gh8	hypothetical gene ZD52F10	727.4433	605	223.8014		-1.70E+00		2 Unchanged Medium
	Q9Y2S6	D01ef2	hypothetical protein	1188,199	1490	1593.015	1423.596	4.23E-01		2 Unchanged Medium
	Q9UKZ1	123gh3	hypothetical protein C40	657.2427	614	506.0272	592,3978	-3.77E-01		2 Unchanged Medium
	290779	D21gh8	hypothetical protein CAB56184	432.491	695	601.25	576,3525	4.75E-01	6.85E-01	2 Unchanged Medium
	095891	D08ef8	hypothetical protein CL25022	1675.198	2070	1477.523	1741.841	-1.81E-01	3.07E-01	2 Unchanged Medium
	2911119	F08gh6	hypothetical protein DJ328E19.C	1119.481	935	1005.691		-1.55E-01		2 Unchanged Medium
	29H0U3	K20gh8	hypothetical protein DKFZp564K	1287.049	1080	1145.302		-1.68E-01		2 Unchanged Medium
	29NWD8		hypothetical protein FLJ10099	754.9838	758		741.1753		5.11E-03	2 Unchanged Medium
	29NW90	P21gh2	hypothetical protein FLJ10211	492.5962	551		548,3816		1.61E-01	2 Unchanged Medium
	29NW16	B18gh2	hypothetical protein FLJ10297	772.0358	844		745.7235		1.29E-01	2 Unchanged Medium
	29NVZ3	G04gh3 G14gh3	hypothetical protein FLJ10374	756.2846	543	467,9722		-6.93E-01		2 Unchanged Medium
	29NVC3	D23gh3	hypothetical protein FLJ10420	437.615	702	362.923	500.7932		6,81E-01	2 Unchanged Medium
	29H9K7	B10gh5	hypothetical protein FLJ10815 hypothetical protein FLJ12681	1402,842	928	598.595		1.23E+00		2 Unchanged Medium
	29H8H4	H06gh5	hypothetical protein FLJ12800	446.1454	657	593.5606	565.6072		5.59E-01	2 Unchanged Medium
	29P0R5	L02ef1	hypothetical protein FLJ14868	1058.023 702,1841	775 684	1421.894	1084.852	4.26E-01		2 Unchanged Medium
	29NXQ4	M07gh3	hypothetical protein FLJ20113	1503,987		1117.168	834.4521 1521.653	6.70E-01		2 Unchanged Medium
	29NXI2	M21gh3	hypothetical protein FLJ20234	643.4985	877	1545.141 178.0291	566,3149		1.13E-02	2 Unchanged Medium
	29NXD5	E18gh2	hypothetical protein FLJ20309	617.1482		758.0334	932,5386	2.97E-01	4.47E-01	2 Unchanged Medium
		116gh2	hypothetical protein FLJ20419	938.8958			915.2399	6.42E-02		2 Unchanged Medium
	29NWX1		hypothetical protein FLJ20552	1366,306					3.95E-01	2 Unchanged Medium
	BTWNE		hypothetical protein FLJ20508	980.6383			934,2965			2 Unchanged Medium
	SWM/N6		hypothetical protein FLJ20739	381,202	593	689.893	554,671		6.37E-01	2 Unchanged Medium
(9H7C7	D05gh5	hypothetical protein FLJ21044 si			585.1487	615 0164	8.53E-02		2 Unchanged Medium 2 Unchanged Medium
(19H6R6	J23gh5		425.3152				6.75E-01		2 Unchanged Medium
		F11gh5	hypothetical protein FLJ22548 sl					1.39E+00		2 Unchanged Medium
		H15ef8		1310.097				1.81E-01		2 Unchanged Medium
	19Y244	B23ef2	hypothetical protein HSPC014	1668.101				-4.23E-01		2 Unchanged Medium
		P04ef2		666.1765	521			4.97E-02 ·		2 Unchanged Medium
		P06ef2		916.5214	1450	1133.279			6.63E-01	2 Unchanged Medium
		P14ef2	hypothetical protein HSPC155	1031.658	1330	662,2481	1008.218		3.67E-01	2 Unchanged Medium
		A19ef3	hypothetical protein HSPC 194	1336.172		1957.055	1622.151		2,36E-01	2 Unchanged Medium
		A23ef3		280.0356	644		532.0086	1.26E+00	1.20E+00	2 Unchanged Medium
		C18ef2		777.5591		551.8354	679,3596			2 Unchanged Medium
		K02ef2		1224.056		1457.544	1322,295	2.52E-01	7.04E-02	2 Unchanged Medium
		K10ef2		376.8259		1014.441		1.43E+00	1.31E+00	2 Unchanged Medium
		K20ef2 K22ef2		1301.286				5.49E-01		2 Unchanged Medium
			hypothetical protein LOC51256	540,326				1.61E-01 -		2 Unchanged Medium
		M23ef2		600.1582					2.52E-01	2 Unchanged Medium
		M02gh4 L24gh7		663.8685 560.8856				3.63E-02 ·		2 Unchanged Medium
		C09gh6		569.8856 1446 395				3.63E-01 ·		2 Unchanged Medium
		K05gh6		1446.395 527.9287			1754.667		7.68E-01	2 Unchanged Medium
				JE1.JE01	619	677.2851	628.1009	3.59E-01	3.63E-01	2 Unchanged Medium

889 899.9017 876.8925 9.67E-02 7.95E-02

2 Unchanged Medium

Q9BQD3 E17gh6 hypothetical protein MGC2749 841.5499

	E1/gh6	hypothetical protein MGC2749	4224.005	889	899,9017	676.8925	9.675-02	7.95E-02	2 Unchanged Medium
Q98Q61	A09gh6	hypothetical protein MGC2803	1234.395	2170	1980.421	1794.728	6.82E-01		2 Unchanged Medium
Q9BTV4	O23gh6	hypothetical protein MGC3222	561.6572	502	583.5781	548.9637	5.52E-02		2 Unchanged Medium
Q9BVX2	C17gh6	hypothetical protein MGC5576	747.2749	762	604.3725	704.7101		2.91E-02	2 Unchanged Medium
Q9H773	117gh6	hypothetical protein MGC5627	799.182	819	584.0079		-4.53E-01	3.57E-02	2 Unchanged Medium
Q9BPX5	LD2gh7	hypothetical protein similar to act		424	632.4101	546.6359	1.16E-01		2 Unchanged Medium
Q9UJX8	B09ef3	hypothetical protein; estradiol-inc		948	1116.899	1083,179	-8.46E-02	-3,21E-01	2 Unchanged Medium
P00492	P12cd6	hypoxanthine phosphoribosyltrar	1210.895	233	566.0945	669,9437	-1.10E+00	-2.38E+00	2 Unchanged Medium
P22304	104ab6	Iduronate 2-sulfatase (Hunter syr	1197.3	1320	1095.895	1205.085	-1.28E-01	1.43E-01	2 Unchanged Medium
Q03827	F02ef6	Immediate early protein	650.6358	537	536,2705	574,4895	-2.79E-01	-2.78E-01	2 Unchanged Medium
P12268	F11ab6	IMP (inosine monophosphate) de	1254.916	567	1293,693	1038.44		-1.15E+00	2 Unchanged Medium
Q9Y6K9	P03cd3	inhibitor of kappa light polypeptic		788	807,6888		-2.60E-02		2 Unchanged Medium
Q14573	F04ab6	inositol 1;4;5-triphosphate recept		646	627.6143	609,9521	1.75E-01	2.17E-01	2 Unchanged Medium
015357	L09ab8	inositol polyphosphate phosphati		1150	1290.473	1146.828	3.71E-01	2.08E-01	2 Unchanged Medium
015503	B05ab7	insulin induced gene 1	472.0202	428	615.7553	505,2037	3.84E-01		
P52945			1437.285	3700					
	L13gh6	Insulin upstream factor 1	1227,459	1290	746.514		-9.45E-01		2 Unchanged Medium
P51460	807ab7	insulin-like 3 (Leydig cell)			1040.851		-2.38E-01	6.76E-02	2 Unchanged Medium
P46977	D20ab6	Integral membrane protein 1	1693.474	1790	2514.977	1998.711	5.71E-01	7.81E-02	2 Unchanged Medium
Q13418	H08ef5	integrin-linked kinase	1340.442	. 962	1599.336	1300,502		-4.79E-01	2 Unchanged Medium
P48551	O21ef7		605.1613	669	389.178		-6.37E-01	1.44E-01	2 Unchanged Medium
P15260	A02ef7	Interferon gamma receptor 1	1165.784	2600	1179.083	1648.504		1.16E+00	2 Unchanged Medium
P38484	D12ef5	interferon gamma receptor 2 (inte	593.8133	773	550.2678	638.9669	-1.10E-01	3.80E-01	2 Unchanged Medium
O14896	K05ef6	Interferon regulatory factor 6	495.9782	951	406.9438	617.9736	-2.85E-01	9.39E-01	2 Unchanged Medium
	G13ef5	interferon-gamma receptor (IFNG	616.4461	474	457.8757	515,9592	-4.29E-01	-3,80E-01	2 Unchanged Medium
Q14116	L03ab6	interleukin 18 (interferon-gamma	1525.444	1610	556.9614	1230,533	-1.45E+00	7.71E-02	2 Unchanged Medium
075874	H11ab7	isocitrate dehydrogenase 1 (NAL	660.9773	496	904.897	687,2716	4.53E-01	-4.14E-01	2 Unchanged Medium
P51553	121gh1	Isocitrate dehydrogenase 3 (NAC		899	784.1063	632,7908	-5.67E-02		2 Unchanged Medium
P41252	P23ab6	Isoleucine-IRNA synthetase	643.6728	316	992.267	650,6946		-1.03E+00	2 Unchanged Medium
Q9UE99	J07ef7	jagged 2	657.9935	460	696,5019	604,7722	8.21E-02		2 Unchanged Medium
P52292	J14ab6	karyopherin alpha 2 (RAG cohori		1270	1670.406	1350.126	5.95E-01		2 Unchanged Medium
P24390	E07cd7	KDEL (Lys-Asp-Glu-Leu) endopl:		1210	1411.383		-1.41E-01		
P08779	D03ab7	keratin 16 (focal non-epidermoly)		786	663.0228		-2.99E-01		
Q04695		keratin 17		1180					
Q04693 Q07666	O24ab6	KH domain containing; RNA bind	580,3123		1056.274	937,9159		1.02E+00	2 Unchanged Medium
	B04cd6			780	1103.092	1031.926		-6.37E-01	2 Unchanged Medium
Q14165	O16gh1	KIAA0152 gene product	285.0232	348	1374.81	669.3755	2.27E+00	2.89E-01	2 Unchanged Medium
Q12765	F03gh1	KIAA0193 gene product	951.658	1000	981.6527	979,0013	4.48E-02	7.68E-02	2 Unchanged Medium
043310	F11gh1	KIAA0427 gene product	482,1741	589	556.0999	542,3144	2.06E-01	2.88E-01	2 Unchanged Medium
043167	J03gh1	KIAA0441 gene product	1398.244	982	1102.589		-3.43E-01		2 Unchanged Medium
Q9H7D5	118gh7	KIAA1191 protein	886,4026	1930	1061.302	1291.483		1.12E+00	2 Unchanged Medium
Q9HD96	L03cd8	LAG1 longevity assurance homol		798	1016,572		-3.76E-01		2 Unchanged Medium
000182	P04ab6	lectin; galactoside-binding; solub		1300	527.6081		-5.28E-01	7.79E-01	2 Unchanged Medium
Q99538	B04ef7	legumain	464.5	517	1105.037	695,3901	1.25E+00	1.53E-01	2 Unchanged Medium
O95751	G01cd8	leucine zipper, down-regulated ir	874.8344	274	970.9239	706.7407	1.50E-01	-1.67E+00	2 Unchanged Medium
O15468	P03ef4	leukocyte immunoglobulin-like re	826.8073	939	1120.204	962.0679	4.38E-01	1.84E-01	2 Unchanged Medium
Q9UBB4	106cd8	like mouse brain protein E46	877.3997	623	692.5911	731,0054	-3.41E-01	-4.94E-01	2 Unchanged Medium
Q9UFW4	N17ef8	likely ortholog of rat golgi stackin	1680.361	1700	2121.05	1834.238	3.36E-01	1.79E-02	2 Unchanged Medium
Q14847	P09ab7	LIM and SH3 protein 1	664.6143	758	895.1972	772.769	4.30E-01	1.91E-01	2 Unchanged Medium
Q00158	M09ef3	LIM domain only 4	457.4088	299	1353,355	703.0924		-6.16E-01	2 Unchanged Medium
Q9HAP6	M16gh5	Lin-7b protein; likely ortholog of a	2114.229	1550	2175,236	1945,862		-4.50E-01	2 Unchanged Medium
P38571	K08ab6	lipase A; lysosomal acid; cholest-		587	512.8879		-2.36E-01		2 Unchanged Medium
Q99732	118cd5			893	449,4711	674.31	-5.98E-01		
Q9Y4Z1		LPS-induced TNF-alpha factor	680.3876						2 Unchanged Medium
		LPS-Induced TNF-elpha factor Lsm3 protein	680.3876 1641.224	1300	1596.905			3.92E-01 -3.40F-01	2 Unchanged Medium 2 Unchanged Medium
075896	O21ef4	Lsm3 protein	1641.224	1300	1596.905	1511.517	-3.95E-02	-3,40E-01	2 Unchanged Medium
O75896 O06643	O21ef4 D05cd7	Lsm3 protein lung cancer candidate	1641.224 690.6061	772	601.3296	1511.517 688.0098	-3.95E-02 -2.00E-01	-3.40E-01 1.61E-01	2 Unchanged Medium 2 Unchanged Medium
Q06643	O21ef4 D05cd7 O07ef6	Lsm3 protein lung cancer candidate lymphotoxin beta (TNF superfam	1641.224 690.6061 405.7368	772 1040	601.3296 298.9254	1511.517 688.0098 582.7217	-3.95E-02 -2.00E-01 -4.41E-01	-3.40E-01 1.61E-01 1.36E+00	2 Unchanged Medium 2 Unchanged Medium 2 Unchanged Medium
Q06643 O95372	O21ef4 D05cd7 O07ef6 J04ef7	Lsm3 protein lung cancer candidate lymphotoxin beta (TNF superfam lysophospholipase ti	1641.224 690.6061 405.7368 480.0562	772 1040 737	601.3296 298.9254 752.7079	1511.517 688.0098 582.7217 656,485	-3.95E-02 -2.00E-01 -4.41E-01 6,49E-01	-3.40E-01 1.61E-01 1.36E+00 6.18E-01	2 Unchanged Medium 2 Unchanged Medium 2 Unchanged Medium 2 Unchanged Medium
Q06643 O95372 P13473	O21ef4 D05cd7 O07ef6 J04ef7 N06ab6	Lsm3 protein lung cancer candidate lymphotoxin beta (TNF superfam lysophospholipase II lysosomal-associated membrane	1641.224 690.6061 405.7368 480.0562 952.6393	772 1040 737 2000	601.3296 298.9254 752.7079 1277.773	1511.517 688.0098 582.7217 656,485 1408.92	-3.95E-02 -2.00E-01 -4.41E-01 6.49E-01 4.24E-01	-3.40E-01 1.61E-01 1.36E+00 6.18E-01 1.07E+00	2 Unchanged Medium 2 Unchanged Medium 2 Unchanged Medium 2 Unchanged Medium 2 Unchanged Medium
Q06643 O95372 P13473 Q08397	O21ef4 D05cd7 O07ef6 J04ef7 N06ab6 F09ab7	Lsm3 protein lung cancer candidate lymphotoxin beta (TNF superfam lysophospholipase II lysosomal-associated membrane lysyl oxidase-like 1	1641.224 690.6061 405.7368 480.0562 952.6393 1391.453	772 1040 737 2000 1630	601.3296 298.9254 752.7079 1277.773 2243.026	1511.517 688.0098 582.7217 656.485 1408.92 1753.531	-3.95E-02 -2.00E-01 -4.41E-01 6.49E-01 4.24E-01 6.89E-01	-3.40E-01 1.61E-01 1.36E+00 6.18E-01 1.07E+00 2.25E-01	2 Unchanged Medium
Q06643 O95372 P13473 Q08397 Q15046	O21ef4 D05cd7 O07ef6 J04ef7 N06ab6 F09ab7 B17ab7	Lsm3 protein lung cancer candidate lymphotoxin beta (TNF superfam lysophospholipase II lysosomal-associated membrane lysyl oxidase-like 1 lysyl-tiRNA synthetase	1641.224 690.6061 405.7368 480.0562 952.6393 1391.453 900.8177	772 1040 737 2000 1630 556	601.3296 298.9254 752.7079 1277.773 2243.025 539.813	1511.517 688.0098 582.7217 656.485 1408.92 1753.531 665.5485	-3.95E-02 -2.00E-01 -4.41E-01 6.49E-01 4.24E-01 6.89E-01 -7.39E-01	-3.40E-01 1.61E-01 1.36E+00 6.18E-01 1.07E+00 2.25E-01 -6.96E-01	2 Unchanged Medium 2 Unchanged Medium
Q06643 O95372 P13473 Q08397 Q15046 Q13312	O21ef4 D05cd7 O07ef6 J04ef7 N06ab6 F09ab7 B17ab7 D17cd3	Lsm3 protein lung cancer candidate lymphotoxin beta (TNF superfam lysophospholipase II lysosomal-associated membrane lysyl oxidase-like t lysyl-IRNA synthetase MAD1 mitotic arrest deficient-like	1641.224 690.6061 405.7368 480.0562 952.6393 1391.453 900.8177 807.7267	772 1040 737 2000 1630 556 503	601.3296 298.9254 752.7079 1277.773 2243.025 539.813 574.4266	1511.517 688.0098 582.7217 656,485 1408.92 1753.531 665,5485 628.231	-3.95E-02 -2.00E-01 -4.41E-01 6.49E-01 4.24E-01 6.89E-01 -7.39E-01 -4.92E-01	-3.40E-01 1.61E-01 1.36E+00 6.18E-01 1.07E+00 2.25E-01 -6.96E-01 -6.85E-01	2 Unchanged Medium
Q06643 O95372 P13473 Q08397 Q15046 Q13312 Q09160	O21ef4 D05cd7 O07ef6 J04ef7 N06ab6 F09ab7 B17ab7 D17cd3 A05ab6	Lsm3 protein lung cancer candidate lymphotoxin beta (TNF superfam lysophospholipase II lysosomal-associated membrane lysyl oxidase-like 1 lysyl-IRNA synthetase MAD1 mitotic arrest deficient-like major histocompatibility complex;	1641.224 690.6061 405.7368 480.0562 952.6393 1391.453 900.8177 807.7267 1524.076	772 1040 737 2000 1630 556 503 2710	601.3296 298.9254 752.7079 1277.773 2243.025 539.813 574.4266 1290.763	1511.517 688.0098 582.7217 656,485 1408.92 1753.531 665.5485 628.231 1841.106	-3.95E-02 -2.00E-01 -4.41E-01 6.49E-01 4.24E-01 6.89E-01 -7.39E-01 -4.92E-01 -2.40E-01	-3.40E-01 1,61E-01 1.36E+00 6.18E-01 1.07E+00 2.25E-01 -6.96E-01 8.30E-01	2 Unchanged Medium
Q06643 O95372 P13473 Q08397 Q15046 Q13312 Q09160 P40925	O21ef4 D05cd7 O07ef6 J04ef7 N06ab6 F09ab7 B17ab7 D17cd3 A05ab6 J23ab7	Lsm3 protein lung cancer candidate lymphotoxin beta (TNF superfam lysophospholipase II lysosomal-associated membrane lysyl oxidase-like 1 lysyl-IRNA synthetase MAD1 mitotic arrest deficient-like major histocompatibility complex; malate dehydrogenase 1; NAD (s	1641.224 690.6061 405.7368 480.0562 952.6393 1391.453 900.8177 807.7267 1524.076 1144.218	772 1040 737 2000 1630 556 503 2710 788	601.3296 298.9254 752.7079 1277.773 2243.025 539.813 574.4266 1290.763 1089.134	1511.517 688.0098 582.7217 656,485 1408.92 1753.531 665,5485 628.231 1841.106	-3.95E-02 -2.00E-01 -4.41E-01 6.49E-01 4.24E-01 6.89E-01 -7.39E-01 -4.92E-01 -2.40E-01 -7.12E-02	-3.40E-01 1.61E-01 1.36E+00 6.18E-01 1.07E+00 2.25E-01 -6.96E-01 -6.85E-01 8.30E-01 -5.38E-01	2 Unchanged Medium
Q06643 O95372 P13473 Q08397 Q15046 Q13312 Q09160 P40925 P40926	O21ef4 D05cd7 O07ef6 J04ef7 N06ab6 F09ab7 B17ab7 D17cd3 A05ab6 J23ab7 L01ab7	Lsm3 protein lung cancer candidate lymphotoxin beta (TNF superfam lysophospholipase II lysosomal-associated membrane lysyl oxidase-like 1 lysyl-IRNA synthetase MAD1 mitotic arrest deficient-like major histocompatibility complex; malate dehydrogenase 1; NAD (s malate dehydrogenase 2; NAD (r	1641.224 690.6061 405.7368 480.0562 952.6393 1391.453 900.8177 807.7267 1524.076 1144.218 1525.837	772 1040 737 2000 1630 556 503 2710 788 1020	601.3296 298.9254 752.7079 1277.773 2243.025 539.813 574.4266 1290.763 1089.134 1212.544	1511.517 688.0098 582.7217 656,485 1408.92 1753.531 665,5485 628.231 1841.106 1007.157 1251.158	-3.95E-02 -2.00E-01 -4.41E-01 6.49E-01 4.24E-01 -7.39E-01 -4.92E-01 -2.40E-01 -7.12E-02 -3.32E-01	-3.40E-01 1.61E-01 1.36E+00 6.18E-01 1.07E+00 2.25E-01 -6.96E-01 -6.85E-01 8.30E-01 -5.38E-01	2 Unchanged Medium
Q06643 Q95372 P13473 Q08397 Q15046 Q13312 Q09160 P40925 P40926 Q16626	O21ef4 D05cd7 O07ef6 J04ef7 N06ab6 F09ab7 B17ab7 D17cd3 A05ab6 J23ab7 L01ab7 J15ef3	Lsm3 protein lung cancer candidate lymphotoxin beta (TNF superfam lysophospholipase II lysosomal-associated membrane lysyl oxidase-like 1 lystyl-IRNA synthetase MAD1 mitotic arrest deficient-like raigir histocompatibility complex; malate dehydrogenase 1; NAD (s mala-enhanced antigen	1641.224 690.6061 405.7368 480.0562 952.6393 1391.453 900.8177 807.7267 1524.076 1144.218 1525.637 1253.21	772 1040 737 2000 1630 556 503 2710 788 1020 1530	601.3296 298.9254 752.7079 1277.773 2243.025 539.813 574.4266 1290.763 1089.134 1212.544 1827.301	1511.517 688.0098 582.7217 656.485 1408.92 1753.531 665.5485 628.231 1841.106 1007.157 1251.158 1536.52	-3.95E-02 -2.00E-01 -4.41E-01 6.49E-01 4.24E-01 6.89E-01 -7.39E-01 -4.92E-01 -2.40E-01 -7.12E-02 -3.32E-01 5.44E-01	-3.40E-01 1.61E-01 1.36E+00 6.18E-01 1.07E+00 2.25E-01 -6.96E-01 -6.85E-01 8.30E-01 -5.38E-01 2.87E-01	2 Unchanged Medium
Q06643 O95372 P13473 Q08397 Q15046 Q13312 Q09160 P40925 P40926 Q16626 Q9NQG1	O21ef4 D05cd7 O07ef6 J04ef7 N06ab6 F09ab7 B17ab7 D17cd3 A05ab6 J23ab7 L01ab7 J15ef3 C06gh5	Lsm3 protein lung cancer candidate lymphotoxin beta (TNF superfam lysophospholipase II lysosomal-associated membrane lysyl oxidase-like 1 lysyl-tRNA synthetase MAD1 mitotic arrest deficient-like major histocompatibility complex; malate dehydrogenase 1; NAD (s malate dehydrogenase 2; NAD (male-enhanced antigen marmosidase; beta A: lysosomal-	1641.224 690.6061 405.7368 480.0562 952.6393 1391.453 900.8177 807.7267 1524.076 1144.218 1525.837 1253.21 533.2883	772 1040 737 2000 1630 556 503 2710 788 1020 1530 800	601.3296 298.9254 752.7079 1277.773 2243.025 539.813 574.4266 1290.783 1089.134 1212.544 1827.301 736.627	1511.517 688.0098 582.7217 656.485 1408.92 1753.531 665.5485 628.231 1841.106 1007.157 1251.158 1536.52 689.9075	-3.95E-02 -2.00E-01 -4.41E-01 6.49E-01 4.24E-01 6.89E-01 -7.39E-01 -2.40E-01 -7.12E-02 -3.32E-01 5.44E-01 4.66E-01	-3.40E-01 1.61E-01 1.36E+00 6.18E-01 1.07E+00 2.25E-01 -6.96E-01 -6.95E-01 5.30E-01 -5.38E-01 2.97E-01 5.85E-01	2 Unchanged Medium
Q06643 Q95372 P13473 Q08397 Q15046 Q13312 Q09160 P40925 P40926 Q16626 Q9NQG1 P26572	O21ef4 D05cd7 O07ef6 J04ef7 N06ab6 F09ab7 B17ab7 D17cd3 A05ab6 J23ab7 L01ab7 J15ef3 C06gh5 G11ab7	Lsm3 protein lung cancer candidate lymphotoxin beta (TNF superfam lysophospholipase II lysosomal-associated membrane lysyl oxidase-like 1 lysyl-IRNA synthetase MAD1 mitotic arrest deficient-like rrajor histocompatibility complex; malate dehydrogenase 1; NAD (s mala-enhanced antigen marnosidase; beta A: lysosomal-mannosyl (alpha-1;3-)-glycoprote	1641.224 690.6061 405.7368 480.0562 952.6393 1391.453 900.8177 807.7267 1524.076 1144.218 1525.837 1253.21 533.2883 619.6733	772 1040 737 2000 1630 556 503 2710 768 1020 1530 800 369	601.3296 298.9254 752.7079 1277.773 2243.026 539.813 574.4266 1290.763 1089.134 1212.544 1827.301 736.627 887.6535	1511.517 688.0098 582.7217 656.485 1408.92 1753.531 665.5485 628.231 1841.106 1007.157 1251.158 1536.52 689.9075 625.3732	-3.95E-02 -2.00E-01 -4.41E-01 6.49E-01 4.24E-01 6.89E-01 -7.39E-01 -2.40E-01 -7.12E-02 -3.32E-01 5.48E-01 5.18E-01	-3.40E-01 1.61E-01 1.36E+00 6.18E-01 1.07E+00 2.25E-01 -6.85E-01 8.30E-01 -5.38E-01 2.87E-01 5.85E-01 -7.49E-01	2 Unchanged Medium
Q06643 O95372 P13473 Q08397 Q15046 Q13312 Q09160 P40925 P40926 Q16626 Q9NQG1 P26572 Q10469	O21ef4 D05cd7 O07ef6 J04ef7 N06ab6 F09ab7 B17ab7 D17cd3 A05ab6 J23ab7 L01ab7 J15ef3 C06gh5 G11ab7 G15ab7	Lsm3 protein lung cancer candidate lymphotoxin beta (TNF superfam lysophospholipase II lysosomal-associated membrane lysyl oxidase-like 1 lystyl-IRNA synthetase MAD1 mitotic arrest deficient-like raigir histocompatibility complex; malate dehydrogenase 1; NAD (s mala-enhanced antigen marmosidase; beta At lysosomal-mannosyl (alpha-1;3-)-glycoprote mannosyl (alpha-1;3-)-glycoprote mannosyl (alpha-1;5-)-glycoprote	1641.224 690.6061 405.7368 480.0562 952.6393 1391.453 900.8177 807.7267 1524.076 1144.218 1525.837 1253.21 533.2883 619.6733 600.1049	772 1040 737 2000 1630 556 503 2710 788 1020 1530 800 369 583	601.3296 298.9254 752.7079 1277.773 2243.026 539.813 574.4266 1290.763 1089.134 1212.544 1827.301 736.627 887.6535	1511.517 688.0098 582.7217 656.485 1408.92 1753.531 665.5485 628.231 1841.106 1007.157 1251.158 1536.52 689.9075	-3.95E-02 -2.00E-01 -4.41E-01 6.49E-01 4.24E-01 6.89E-01 -7.39E-01 -2.40E-01 -7.12E-02 -3.32E-01 5.44E-01 4.66E-01 8.52E-02	-3.40E-01 1.61E-01 1.36E+00 6.18E-01 1.07E+00 2.25E-01 -6.96E-01 -8.30E-01 -5.38E-01 2.97E-01 5.88E-01 2.97E-01 5.85E-01 4.29E-02	2 Unchanged Medium
Q06643 Q95372 P13473 Q08397 Q15046 Q13312 Q09160 P40925 P40926 Q16626 Q9NQG1 P26572 Q10469 P41223	O21ef4 D05cd7 O07ef6 J04ef7 N06ab6 F09ab7 B17ab7 D17cd3 A05ab6 J23ab7 L01ab7 J15ef3 C06gh5 G11ab7 L09cd4	Lsm3 protein lung cancer candidate lymphotoxin beta (TNF superfam lysophospholipase II lysosomal-assodated membrane lysyl oxidase-like 1 lysyl-tRNA synthetase MAD1 mitotic arrest deficient-like major histocompatibility complex; malate dehydrogenase 1; NAD (s malate dehydrogenase 2; NAD) (s m	1641.224 690.6061 405.7368 480.0562 952.6393 1391.453 900.8177 807.7267 1524.076 1144.218 1253.21 533.2883 619.6733 600.1049 372.2443	772 1040 737 2000 1630 556 503 2710 768 1020 1530 800 369	601.3296 298.9254 752.7079 1277.773 2243.025 539.813 574.4266 1290.763 1089.134 1212.544 1827.301 736.627 887.6535 636.621	1511.517 688.0098 582.7217 656.485 1408.92 1753.531 665.5485 628.231 1841.106 1007.157 1251.158 1536.52 689.9075 625.3732 606.4146 5003.5179	-3.95E-02 -2.00E-01 -4.41E-01 6.49E-01 4.24E-01 6.89E-01 -7.39E-01 -4.92E-01 -7.12E-02 -3.32E-01 5.44E-01 4.66E-01 5.18E-01 6.52E-02 6.29E-01	-3.40E-01 1.51E-01 1.36E+00 6.18E-01 1.07E+00 2.25E-01 6.96E-01 -6.96E-01 6.30E-01 5.38E-01 2.97E-01 5.85E-01 7.49E-01 5.96E-01 5.96E-01	2 Unchanged Medium
Q06643 O95372 P13473 Q08397 Q15046 Q13312 Q09160 P40925 P40926 Q16626 Q9NQG1 P26572 Q10469	O21ef4 D05cd7 O07ef6 J04ef7 N06ab6 F09ab7 B17ab7 D17cd3 A05ab6 J23ab7 L01ab7 J15ef3 C06gh5 G11ab7 G15ab7	Lsm3 protein lung cancer candidate lymphotoxin beta (TNF superfam lysophospholipase II lysosomal-associated membrane lysyl oxidase-like 1 lystyl-IRNA synthetase MAD1 mitotic arrest deficient-like raigir histocompatibility complex; malate dehydrogenase 1; NAD (s mala-enhanced antigen marmosidase; beta At lysosomal-mannosyl (alpha-1;3-)-glycoprote mannosyl (alpha-1;3-)-glycoprote mannosyl (alpha-1;5-)-glycoprote	1641.224 690.6061 405.7368 480.0562 952.6393 1391.453 900.8177 807.7267 1524.076 1144.218 1525.837 1253.21 533.2883 619.6733 600.1049	772 1040 737 2000 1630 556 503 2710 788 1020 1530 800 369 583	601.3296 298.9254 752.7079 1277.773 2243.025 539.813 574.4266 1290.763 1089.134 1212.544 1827.301 736.627 887.6535 636.621	1511.517 688.0098 582.7217 656.485 1408.92 1753.531 665.5485 628.231 1841.106 1007.157 1251.158 1536.52 689.9075 625.3732 606.4146 5003.5179	-3.95E-02 -2.00E-01 -4.41E-01 6.49E-01 4.24E-01 6.89E-01 -7.39E-01 -2.40E-01 -7.12E-02 -3.32E-01 5.44E-01 4.66E-01 8.52E-02	-3.40E-01 1.51E-01 1.36E+00 6.18E-01 1.07E+00 2.25E-01 6.96E-01 -6.96E-01 6.30E-01 5.38E-01 2.97E-01 5.85E-01 7.49E-01 5.96E-01 5.96E-01	2 Unchanged Medium
Q06643 Q95372 P13473 Q08397 Q15046 Q13312 Q09160 P40925 P40926 Q16626 Q9NQG1 P26572 Q10469 P41223	O21ef4 D05cd7 O07ef6 J04ef7 N06ab6 F09ab7 B17ab7 D17cd3 A05ab6 J23ab7 L01ab7 J15ef3 C06gh5 G11ab7 L09cd4	Lsm3 protein lung cancer candidate lymphotoxin beta (TNF superfam lysophospholipase II lysosomal-assodated membrane lysyl oxidase-like 1 lysyl-tRNA synthetase MAD1 mitotic arrest deficient-like major histocompatibility complex; malate dehydrogenase 1; NAD (s malate dehydrogenase 2; NAD) (s m	1641.224 690.6061 405.7368 480.0562 952.6393 1391.453 900.8177 807.7267 1524.076 1144.218 1525.837 1253.21 533.2883 619.6733 600.1043 372.2443 397.4495	772 1040 737 2000 1630 556 503 2710 788 1020 1530 800 369 583 563	601.3296 298,9254 752.7079 1277.773 2243.025 539.813 574.4266 1290.763 1089,134 1212.544 1827.301 736.627 887.6535 636.621 575.7695	1511.517 688.0098 582.7217 656.485 1408.92 1753.531 665.5485 628.231 1841.106 1007.157 1251.158 1536.52 689.9075 625.3732 606.4146 503.5179 1008.268	-3.95E-02 -2.00E-01 -4.41E-01 6.49E-01 4.24E-01 6.89E-01 -7.39E-01 -4.92E-01 -7.12E-02 -3.32E-01 5.44E-01 4.66E-01 5.18E-01 6.52E-02 6.29E-01	-3.40E-01 1.61E-01 1.36E+00 6.18E-01 1.07E+00 2.25E-01 -6.96E-01 6.36E-01 5.38E-01 2.87E-01 5.85E-01 7.49E-01 4.29E-02 5.96E-01 1.26E+00	2 Unchanged Medium
Q06643 Q95372 P13473 Q08397 Q15046 Q13312 Q09160 P40925 P40926 Q16626 Q9NQG1 P26572 Q10469 P41223 Q9Y5V3	O21ef4 D05cd7 O07ef6 J04ef7 N06ab6 F09ab7 B17ab7 D17cd3 A05ab6 J23ab7 L01ab7 L01ab7 G15ab7 G15ab7 G15ab7 G31ab7 G33ab6	Lsm3 protein lung cancer candidate lymphotoxin beta (TNF superfam lysophospholipase II lysosomal-associated membrane lysyl oxidase-like 1 lysyl-tRNA synthetase MAD1 mitotic arrest deficient-like rnajor histocompatibility complex; malate dehydrogenase 1; NAD (s malate dehydrogenase 2; NAD (s malate dehydrogenase 2; NAD (s malate dehydrogenase 2; NAD (s malate dehydrogenase 3; NAD) (s malate dehydrogenase 4; NAD) (s malate dehydrogenase 4; NAD) (s malate dehydrogenase 2; NAD) (s malate dehydrogenase 1; NAD)	1641.224 690.6061 405.7368 480.0562 952.6393 1391.453 900.8177 807.7267 1524.076 1144.218 1525.837 1253.21 533.2883 619.6733 600.1043 372.2443 397.4495	772 1040 737 2000 1630 556 503 2710 788 1020 1530 800 369 563 563 952	601.3296 298.9254 752.707 2277.773 2243.025 539.813 574.4266 1290.783 1089.134 1212.544 1827.301 736.627 887.635 636.621 575.7695 1674.881	1511.517 688.0098 582.7217 656.485 1408.92 1753.531 665.5485 628.231 1841.106 1007.157 1251.158 1536.52 689.9075 629.3732 606.4146 503.5179 1008.268 704.4837	-3,95E-02 -2,00E-01 -4,41E-01 4,24E-01 6,49E-01 4,24E-01 -7,39E-01 -2,40E-01 -7,12E-02 -3,32E-01 5,44E-01 6,56E-01 5,16E-01 2,08E+00	-3.40E-01 1.61E-01 1.36E+00 6.18E-01 1.07E+00 2.25E-01 -6.96E-01 6.36E-01 5.38E-01 2.87E-01 5.85E-01 7.49E-01 4.29E-02 5.96E-01 1.26E+00	2 Unchanged Medium
Q05643 Q95372 P13473 Q08397 Q15046 Q13312 Q09160 P40925 P40925 Q16626 Q16626 Q16626 Q16626 Q16626 Q16627 Q10423 Q975V3 P43121 Q92494 Q14444	O21ef4 D05cd7 O07ef6 J04ef7 N06ab6 F09ab7 B17ab7 D17cd3 A05ab6 J23ab7 L01ab7 J15ef3 C06gh5 G11ab7 G15ab7 L09cd4 G13cd6 I22ef5	Lsm3 protein lung cancer candidate lymphotoxin beta (TNF superfam lysophospholipase II lysosomal-associated membrane lysyl oxidase-like 1 lysyl-IRNA synthetase MAD1 mitotic arrest deficient-like major histocompatibility complex; malate dehydrogenase 1; NAD (s malate dehydrogenase 2; NAD (s malate enhydrogenase 2; NAD (s malate-enhanced antigen marnosidase; beta A: lysosomal-mannosyl (alpha-1;5-)-glycoprote maternal G10 transcript melanoma antigen; family D; 1 melanoma cell adheslon molecul membrane cofactor protein (CD4 membrane component; chromost	1641.224 690.6061 405.7368 480.0562 952.6393 1391.453 900.8177 807.7267 1524.076 1144.218 1525.837 1253.21 533.2883 619.6733 600.1049 372.2443 397.4495 886.0756 973.362 549.4803	772 1040 737 2000 1630 556 503 2710 788 1020 1530 800 369 583 563 952 795	601.3296 298.9254 752.7079 1277.773 2243.025 539.813 574.4266 1290.783 1089.134 1212.544 1827.301 736.627 887.6535 636.621 575.7685 1674.881 432.1718	1511.517 688.0098 552.7217 656.485 1408.92 1753.531 665.5485 628.231 1841.106 1007.157 1251.158 1536.52 689.9075 625.3732 606.4146 503.5179 1008.268 704.4837 952.9144	-3,95E-02 -2,00E-01 -4,41E-01 -6,49E-01 -4,24E-01 -6,89E-01 -2,40E-01 -7,12E-02 -3,32E-01 -5,44E-01 -4,66E-01 -5,18E-01 -6,29E-01 -2,08E+00 -1,04E+00	-3,40E-01 1,61E-01 0,618E-01 1,07E+00 2,25E-01 -6,95E-01 6,85E-01 8,30E-01 -5,38E-01 2,87E-01 5,85E-01 -7,49E-01 4,29E-02 5,96E-01 1,26E+00 -1,56E-01	2 Unchanged Medium
Q06643 Q95372 P13473 Q08397 Q15046 Q13312 Q09160 P40925 P40926 Q16626 Q9NQG1 P26572 Q10469 P41223 Q9Y5V3 Q9Y5V3 Q92494	O21ef4 D05cd7 O07ef6 J04ef7 N06ab6 F09ab7 B17ab7 D17cd3 A05ab6 J23ab7 J15ef3 C06gh5 G11ab7 G15ab7 L09cd4 G13cd6 G22ef5 D11ef1	Lsm3 protein lung cancer candidate lymphotoxin beta (TNF superfam lysophospholipase II lysosomal-associated membrane lysyl oxidase-like 1 lysyl-IRNA synthetase MAD1 mitotic arrest deficient-like major histocompatibility complex; malate dehydrogenase 1; NAD (s malate dehydrogenase 2; NAD (s malate dehydrogenas	1641.224 690.6061 405.7368 480.0562 952.6393 1391.453 900.8177 807.7267 1524.076 1144.218 1525.837 1253.21 533.2883 619.6733 600.1049 372.2443 397.4495 886.0756 973.362 549.4803	772 1040 737 2000 1630 556 503 2710 788 1020 1530 800 369 583 563 952 7955	601.3296 298.9254 752.7079 1277.773 2243.025 539.813 574.4266 1290.783 1089.134 1212.544 1827.301 736.627 887.6535 636.621 575.7695 1674.881 432.1718 834.044	1511.517 688.0098 552.7217 656.485 1408.92 1753.531 665.5485 628.231 1841.106 1007.157 1251.158 1536.52 689.9075 625.3732 606.4146 503.5179 1008.268 704.4837 952.9144	-3.95E-02 -2.00E-01 -4.41E-01 6.49E-01 4.24E-01 -7.39E-01 -7.12E-02 -3.22E-01 5.44E-01 4.66E-01 5.18E-02 6.29E-01 2.08E+00 -7.04E+00 -7.04E+00 -7.23E-01	-3.40E-01 1.61E-01 1.36E+00 6.18E-01 1.07E+00 2.25E-01 -6.95E-01 -6.35E-01 -5.38E-01 2.07E-01 2.07E-01 4.29E-02 5.59E-01 1.26E+00 -1.56E-01 1.11E-01	2 Unchanged Medium
Q05643 Q95372 P13473 Q08397 Q15046 Q13312 Q09160 P40925 P40925 Q16626 Q16626 Q16626 Q16626 Q16626 Q16627 Q10423 Q975V3 P43121 Q92494 Q14444	O21er4 D05cd7 O07er6 J04er7 N06ab6 F09ab7 B17ab7 D17cd3 A05ab6 J23ab7 L01ab7 J15er3 C06gh5 G11ab7 G15ab7 L09cd4 G13cd6 t22er5 D11er1 H15ab7	Lsm3 protein lung cancer candidate lymphotoxin beta (TNF superfam lysophospholipase II lysosomal-associated membrane lysyl oxidase-like 1 lysyl-IRNA synthetase MAD1 mitotic arrest deficient-like major histocompatibility complex; malate dehydrogenase 1; NAD (s malate dehydrogenase 2; NAD (s malate enhydrogenase 2; NAD (s malate-enhanced antigen marnosidase; beta A: lysosomal-mannosyl (alpha-1;5-)-glycoprote maternal G10 transcript melanoma antigen; family D; 1 melanoma cell adheslon molecul membrane cofactor protein (CD4 membrane component; chromost	1641.224 690.6061 405.7368 480.0562 952.6393 1391.453 900.8177 807.7267 1144.216 1525.837 1253.21 533.2883 619.6733 600.1049 372.2443 397.4495 886.0756 973.362 549.4693 749.6930	772 1040 737 2000 1630 556 503 2710 788 1020 1530 800 369 583 952 795 1050 666	601.3296 298.9254 752.7079 1277.773 2243.025 539.813 574.4266 1290.783 1089.134 1212.544 1827.301 786.627 887.6535 636.621 575,7695 1674.881 432.1718 834.044 464.8174 1085.23	1511.517 688.0098 582.7217 656.485 1408.92 1753.531 665.5485 628.231 1841.106 1007.157 1251.158 1536.52 689.9075 625.3732 606.4146 503.5179 1008.268 704.4837 952.9144 560.0583	-3.95E-02 -2.00E-01 -4.41E-01 6.49E-01 4.24E-01 -7.39E-01 -4.92E-01 -7.12E-02 -3.32E-01 5.44E-01 4.66E-01 5.18E-02 6.29E-01 2.02E-01 -2.23E-01 -2.23E-01 5.33E-01	-3.40E-01 1.61E-01 1.06E+00 6.18E-01 1.07E+00 2.25E-01 -6.85E-01 -6.85E-01 -5.88E-01 5.88E-01 5.88E-01 5.88E-01 1.26E-00 1.26E-00 1.16E-00 1.11E-01 2.77E-01	2 Unchanged Medium 3 Unchanged Medium 3 Unchanged Medium 3 Unchanged Medium 4 Unchanged Medium 5 Unchanged Medium 5 Unchanged Medium 6 Unchanged Medium 7 Unchanged Medium 8 Unchanged Medium 8 Unchanged Medium 9 Unchanged Medium
Q06643 Q95372 Q15046 Q13312 Q09160 P40925 P40926 Q16626 Q9NQG1 P26572 Q10469 P41223 Q9Y5V3 Q43121 Q92494 Q14444 Q3NPE2	O21er4 D05cd7 O7er6 J04er7 N06ab6 F09ab7 B17ab7 D17cd3 A05ab6 J23ab7 L01ab7 L01ab7 C15eb7 L09cd4 G13cd6 G13ab7 L2er5 D11er1 H15ab7 O12er2 G07ab7	Lsm3 protein lung cancer candidate lymphotoxin beta (TNF superfam lysophospholipase II lysosomal-associated membrane lysyl oxidase-like 1 lysyl-IRNA synthetase MAD1 mitotic arrest deficient-like raigir histocompatibility complex; relate dehydrogenase 1; NAD (s malae dehydrogenase 2; NAD (malae-enhanced antigen marmosidase; beta At lysosomal-mannosyl (alpha-1;3-)-glycoprote raternal G10 transcript melanoma antigen; family D; 1 melanoma coffactor protein (CD4 membrane component; chromosc mesenchymal stem cell protein C	1641.224 690.6081 405.7368 480.0562 952.6393 1391.453 900.8177 807.7267 1524.076 1144.218 1525.837 1253.21 533.2883 619.6733 600.1049 372.2443 397.4495 886.0756 973.362 549.4803 749.6938 617.581	772 1040 737 2000 1630 556 503 2710 788 1020 1530 800 369 583 563 952 795 1050 666 932	601.3296 298.9254 752.7079 1277.773 2243.025 539.813 574.4266 1290.783 1089.134 1212.544 187.301 736.627 887.6535 636.621 575.7695 1674.881 432.1718 834.044 464.8174 1085.26 537.1735	1511.517 688.0098 582.7217 656.485 1408.92 1753.531 665.5485 628.231 1841.106 1007.157 1251.158 1536.52 689.9075 625.3732 606.4146 503.5179 1008.268 704.4837 952.9144 560.0593 922.5469 533.4668	-3.95E-02 -2.00E-01 -4.41E-01 6.49E-01 4.24E-01 -7.39E-01 -7.12E-02 -3.22E-01 5.44E-01 5.45E-01 6.29E-01 2.08E-00 -2.23E-01 -2.23E-01 -2.41E-01 -3.52E-02 6.29E-01 -2.23E-01 -2.41E-01 -2.23E-01 -2.23E-01 -2.41E-01	-3.40E-01 1.61E-01 1.05E-00 6.18E-01 1.07E-00 2.25E-01 -6.96E-01 -6.96E-01 -5.30E-01 -5.30E-01 5.85E-01 -7.49E-01 -4.29E-02 -1.56E-01 1.26E+00 -1.56E-01 1.11E-01 2.77E-01 -3.74E-01	2 Unchanged Medium
Q06643 Q95372 Q15046 Q15046 Q13312 Q09160 P40925 P40925 Q9NQ61 P26572 Q10469 P41223 Q9Y5V3 P43121 Q92494 Q14444 Q9NPE2 Q9NPE2	021er4 D05cd7 O07er6 J04er7 N06ab6 F09ab7 B17ab7 D17cd3 A05ab6 J23ab7 L01ab7 J15er3 C06gh5 G11ab7 G15ab7 L09cd4 G13cd5 I22er5 D11er1 D12er2 G07ab7 N11er2 N11er2 N11er2 N11er3 N11er3 N11er3 N11er3 N11er3 N11er3	Lsm3 protein lung cancer candidate lymphotoxin beta (TNF superfam lysophospholipase II lysosomal-associated membrane lysyl oxidase-like 1 lysyl-IRNA synthetase MAD1 mitotic arrest deficient-like rajor histocompatibility complex; malate dehydrogenase 1; NAD (s male-enhanced antigen marnosidase; beta A: lysosomal-mannosyl (alpha-1;3-)-glycoprote maternal G10 transcript melanoma antigen; family D; 1 melanoma cell adheslon molecul membrane cofactor protein (CD4 membrane corponent; chromosx mesenchymal stem cell protein C mesoderm specific transcript hon metalloprotease 1 (pliritysin famil	1641.224 690.6081 405.7368 480.0562 952.6393 1391.453 900.8177 807.7267 1524.076 1144.218 1525.837 1253.21 533.2883 619.6733 600.1049 372.2443 397.4495 886.0756 973.362 549.4803 749.6938 617.581	772 1040 737 2000 1630 556 503 2710 768 1020 1530 800 369 583 563 3952 795 1050 666 932 446 446 349	601.3296 298.9254 752.7079 1277.773 2243.025 539.813 574.4266 1290.783 1089.134 1212.544 1827.301 736.627 887.6535 636.621 575.7695 1674.881 432.1718 834.044 464.8174 1085.26 535 478.8153	1511.517 688.0098 582.7217 656.485 1408.92 1753.531 665.5485 628.231 1841.106 1007.157 1251.158 1536.52 689.9075 625.3732 606.4146 503.5179 1008.268 704.4837 952.9144 953.34868 520.889 520.889	-3.95E-02 -2.00E-01 -4.41E-01 -4.41E-01 -4.49E-01 -7.39E-01 -4.92E-01 -2.40E-01 -7.12E-02 -3.32E-01 -5.44E-01 -5.46E-01 -5.18E-01 -1.04E+00 -2.23E-01 -2.08E+00 -2.23E-01 -2.41E-01 -5.33E-01 -2.01E-01	-3.40E-01 1.61E-01 1.36E+00 6.18E-01 1.07E+00 0.225E-01 -6.95E-01 -6.85E-01 -5.38E-01 2.87E-01 2.87E-01 4.29E-02 5.96E-01 1.26E+00 -1.56E-01 1.11E-01 2.77E-01 3.41E-01 -1.08E+00	2 Unchanged Medium 3 Unchanged Medium 4 Unchanged Medium 5 Unchanged Medium 5 Unchanged Medium 6 Unchanged Medium 6 Unchanged Medium 7 Unchanged Medium 8 Unchanged Medium 9 Unchanged Medium
Q06643 Q95372 Q15046 Q13312 Q09160 P40925 P40926 Q16626 Q9NQG1 P26572 Q10469 P41223 Q9Y5V3 P43121 Q92494 Q1444 Q14444 Q1444 Q14444 Q14444 Q1444 Q14444 Q14444 Q14444 Q14444 Q14444 Q1444 Q14444 Q1444	O21er4 D05cd7 O7er6 J04er7 N06ab6 F09ab7 B17ab7 D17cd3 A05ab6 J23ab7 L01ab7 L01ab7 L01ab7 L09cd4 G15ab7 L09cd4 G15ab7 L09cd4 G15ab7 L09cd4 G12cd6 U22er5 D11er1 H15ab7 D12er2 G07ab7 N11er3 H20ab7	Lsm3 protein lung cancer candidate lymphotoxin beta (TNF superfam lysophospholipase II lysosomal-associated membrane lysyl oxidase-like 1 lysyl-IRNA synthetase MAD1 mitotic arrest deficient-like major histocompatibility complex; malate dehydrogenase 1; NAD (s malate dehydrogenase 2; NAD (malate dehydrogenase 2; NAD (malate dehydrogenase 2; NAD (s malate dehydrogenase 1;3-)-glycoprote maternal (310 transcript melanoma antigen; family D; 1 melanoma antigen; family D; 1 melanoma coffactor protein (CD4 membrane component; chromosomesenchymal stem cell protein C mesoderm specific transcript hon metalloprotease 1 (plirilysin famil metallothlonein 1).	1641.224 690.6061 405.7368 480.0562 952.6393 1391.453 900.8177 807.7267 1144.218 1525.837 1253.21 533.2883 619.6733 600.1049 372.2443 397.4495 886.0756 973.362 549.4903 617.581 735.157 589.6327	772 1040 737 2000 1630 556 503 2710 788 1020 1530 800 369 583 563 952 795 1050 666 932 446 349 502	601,3296 298,9254 752,7079 1277,773 2243,025 539,813 574,4266 1290,763 1089,134 1212,544 1827,301 736,627 887,6535 636,621 575,7695 1674,881 432,1718 834,044 464,8174 1085,26 537,1735 478,8153 656,4441	1511.517 688.0098 582.7217 656.485 1408.92 1753.531 665.5485 628.231 1841.106 1007.157 1251.158 1536.52 689.9075 625.3732 606.4146 503.5179 1008.268 704.4837 952.9144 560.683 952.546 520.599 522.546 520.899 522.547	-3.95E-02 -2.00E-01 -4.41E-01 6.49E-01 4.24E-01 -7.39E-01 -4.92E-01 -7.12E-02 -3.32E-01 5.44E-01 4.66E-01 5.18E-01 0.52E-02 6.29E-01 2.23E-01 -2.41E-01 -2.31E-01 -2.41E-01 -1.04E+00 -2.23E-01 -2.01E-01 -6.19E-01	-3.40E-01 1.61E-01 1.06E+00 6.18E-01 1.07E+00 2.25E-01 6.85E-01 -6.85E-01 -5.88E-01 5.88E-01 5.88E-01 5.88E-01 7.49E-01 5.96E-01 1.26E-01 1.11E-01 2.47E-01 1.11E-01 2.47E-01 1.11E-01 2.47E-01 1.11E-01 2.47E-01 1.11E-01 2.47E-01	2 Unchanged Medium
Q06643 Q95372 Q15046 Q13312 Q09160 P40925 P40925 Q9NQ61 Q9NQ61 Q92494 Q14444 Q9NPE2 Q9NPE2 Q9PE2	O21er4 D05cd7 O7cr6 J04er7 N06ab6 F09ab7 B17ab7 D17cd3 A05ab6 J23ab7 L01ab7 L01ab7 C15eb7 L09cd4 G13cd6 G13cd6 G13cd7 H15ab7 O12er2 G07ab7 N11er3 H20ab7 C18er7	Lsm3 protein lung cancer candidate lymphotoxin beta (TNF superfam lysophospholipase II lysosomal-assodated membrane lysyl oxidase-like 1 lysyl-IRNA synthetase MAD1 mitotic arrest deficient-like major histocompatibility complex; malate dehydrogenase 1; NAD (s malate dehydrogenase 2; NAD (s mannosyl (alpha-1;3-)-glycoprote mannosyl (alpha-1;3-)-glycoprote maternal G10 transcript melanoma cell adheslon molecul membrane corponent; chromost mesenchymal stem cell protein C mesoderm specific transcript hon metalloprotease 1 (plirilysin famil metalloprotease 1) (plirilysin famil metallothionein 1). microsomal glutathione S-transfe	1641.224 690.6081 405.7368 480.0562 952.6393 1391.453 900.8177 807.7267 1524.076 1144.218 1525.837 1253.21 533.2883 619.6733 600.1049 372.2443 397.4495 886.0756 973.362 549.4803 749.6938 617.581 735.157 589.6327 527.4698	772 1040 737 2000 1630 556 503 2710 768 1020 1530 800 369 583 952 795 1050 666 932 446 349 502 686	601.3296 298.9254 752.7079 1277.773 2243.025 539.813 574.4266 1290.783 1089.134 1212.544 1827.301 736.627 887.6535 636.821 575.7695 1674.881 432.1718 834.044 464.8174 1085.26 537.1735 478.8153 656.4441 373.9779	1511.517 688.0098 582.7217 656.485 1408.92 1753.531 665.5485 628.231 1841.106 1007.157 1251.158 1536.52 689.9075 625.3732 606.4146 503.5179 1008.268 704.4837 952.5469 533.4868	-3.95E-02 -2.00E-01 -4.41E-01 6.49E-01 4.24E-01 -7.39E-01 -7.12E-02 -3.22E-01 5.18E-01 6.52E-01 2.08E-00 -2.23E-01 -2.14E-01 5.18E-01 6.29E-01 -2.23E-01 -2.41E-01 5.18E-01 6.29E-01 -2.23E-01 -2.41E-01 -2.23E-01 -2.41E-01 -2.41E-01 -2.41E-01 -2.41E-01 -2.41E-01 -2.41E-01 -2.41E-01 -2.41E-01 -2.41E-01 -2.41E-01 -2.41E-01 -4.96E-01	-3.40E-01 1.61E-01 1.05E-00 6.18E-01 1.07E-00 2.25E-01 -6.96E-01 -6.96E-01 -5.30E-01 -5.30E-01 5.85E-01 -7.49E-01 -4.29E-02 5.96E-01 1.26E-00 -1.56E-01 -1.7.4E-01 -1.7.4E-01 -1.7.4E-01 -1.00E+00 -2.27E-01	2 Unchanged Medium 3 Unchanged Medium 4 Unchanged Medium 4 Unchanged Medium 5 Unchanged Medium 5 Unchanged Medium 6 Unchanged Medium 6 Unchanged Medium 7 Unchanged Medium 8 Unchanged Medium 9 Unchanged Medium
Q06643 Q95372 Q15046 Q13312 Q09150 P40925 Q16626 Q16626 Q16626 Q16626 Q19523 Q10469 P41223 Q9573 Q14444 Q9NPE2 Q92571 Q95204 P80297 Q99735 Q99735 Q99735 Q99735 Q99735 Q99735	021er4 D05cd7 D07er6 J04er7 N06ab6 F09ab7 B17ab7 D17cd3 A05ab6 J23ab7 L01ab7 J15er3 C06gh5 G11ab7 G15ab7 L09cd4 G13cd6 I22er5 D11er1 H15ab7 O12er2 G07ab7 O12er2 G07ab7 C14er7	Lsm3 protein lung cancer candidate lymphotoxin beta (TNF superfam lysophospholipase II lysosomal-associated membrane lysol oxidase-like 1 lysty-IRNA synthetase MAD1 mitotic arrest deficient-like major histocompatibility complex; malate dehydrogenase 1; NAD (smalate dehydrogenase 2; NAD (malate dehydrogenase 2; NAD (malate dehydrogenase 2; NAD (smalate dehydrogenase) (alpha-1;6-)-glycoprote maternal G10 transcript melanoma antigen; family D; 1 melanoma component; chromosc mesenchymal stem cell protein CD4 metalloprotease 1 (plitrilysin famil metallothionein 1). microsomal glutathione S-transfe microtubule-associated protein 1	1641.224 690.6061 405.7368 480.0562 952.6393 1391.453 900.8177 807.7267 1524.076 1524.076 1524.076 1525.321 533.283 619.6733 600.1049 372.2443 397.4495 886.0756 973.362 549.4803 749.6938 617.581 735.157 589.6327 527.4698	772 1040 737 2000 1630 556 503 2710 768 1020 1530 800 369 583 563 3952 795 1050 666 932 446 349 502 666 692	601.3296 298.9254 752.7079 1277.773 2243.025 539.813 574.4266 1290.783 1089.134 1212.544 1827.301 736.627 887.6535 636.621 575.7695 1674.881 432.1718 834.044 464.8174 1085.26 537.1735 478.8153 678.4441 373.9779 887.6299	1511.517 688.0098 582.7217 656.485 1408.92 1753.531 665.5485 628.231 1841.106 1007.157 1251.158 1536.52 689.9075 625.3732 606.4146 503.5179 1008.268 704.4837 952.9144 503.550.0563 922.5669 533.4668 520.899 582.747 529.1741 699.0441	-3.95E-02 -2.00E-01 -4.41E-01 6.49E-01 4.24E-01 6.69SE-01 -7.39E-01 -4.92E-01 -7.12E-02 -3.32E-01 5.44E-01 4.66E-01 5.18E-01 8.52E-02 6.29E-01 2.29E-01 1.04E+00 -1.04E+00 -2.23E-01 -2.41E-01 1.55E-01 4.96E-01 1.55E-01 4.96E-01	-3.40E-01 1.61E-01 1.06E+00 6.18EE-01 1.07E+00 6.98EE-01 6.98EE-01 6.98EE-01 2.87E-01 5.98EE-01 2.87E-01 4.29E-02 5.96E-00 1.56E-01 1.11E-01 2.77E-01 3.14E-01 -1.08E+00 -2.32E-01 1.149E+00	2 Unchanged Medium 3 Unchanged Medium 4 Unchanged Medium 4 Unchanged Medium 5 Unchanged Medium 5 Unchanged Medium 6 Unchanged Medium 6 Unchanged Medium 6 Unchanged Medium 7 Unchanged Medium 8 Unchanged Medium 9 Unchanged Medium
Q06643 Q95372 Q15046 Q13312 Q09160 P40925 P40926 Q16626 Q9NQG1 P26572 Q10469 P41223 Q9Y5V3 Q43121 Q92494 Q14444 Q3NP62 Q92571 Q95204 Q95271 Q95204 Q95271 Q95205 Q95271 Q95205 Q95275 Q9	O21er4 D05cd7 O7er6 J04er7 N06ab6 F09ab7 B17ab7 D17cd3 A05ab6 J23ab7 L01ab7 L01ab7 L01ab7 L02cd4 G13cd6 t22er5 D11er1 H15ab7 D12er2 G07ab7 N11er3 D13er7 D13g68 M14er7 D13g68 M14er8	Lsm3 protein lung cancer candidate lymphotoxin beta (TNF superfam lysophospholipase II lysosomal-assodated membrane lysyl oxidase-like 1 lysyl-IRNA synthetase MAD1 mitotic arrest deficient-like major histocompatibility complex; malate dehydrogenase 1; NAD (s malate dehydrogenase 2; NAD (mala-enhanced antigen mamosidase; beta A: lysosomal-mannosyl (alpha-1;3-)-glycoprote maternal G10 transcript melanoma antigen; family D: 1 melanoma collactor protein (CD4 membrane component; chromosomesenchymal stem cell protein C mesoderm specific transcript hon metalloprotease 1 (pliritysin famil metallothionein 11. microsomal glutathione S-transle microtubule-associated protein 1 milochondrial carrier homolog 2	1641.224 690.6081 405.7368 480.0562 952.6393 390.8177 807.7267 11252.078 11253.21 132533.21 13253.21 1	772 1040 737 2000 1630 556 503 2710 788 1020 1530 800 369 583 563 952 7955 1050 666 932 446 349 502 686 692	601,3296 298,9254 752,7079 1277,773 2243,025 539,813 574,4266 1290,763 1089,134 1212,544 1827,301 736,627 887,635 636,621 575,7695 1674,881 432,1718 834,044 464,8174 1085,26 537,1735 478,8153 656,4441 173,9779 887,6299	1511.517 688.0098 582.7217 656.485 1408.92 1753.531 665.5485 628.231 1841.106 1007.157 1251.158 1536.52 689.9075 625.3732 606.4146 503.5179 1008.268 704.4837 952.9144 560.0583 952.546 520.899 527.747 529.1741 699.0447 1892.145	-3,95E-02 -2,00E-01 -4,41E-01 6,49E-01 4,24E-01 -7,39E-01 -4,92E-01 -7,12E-02 -3,32E-01 5,44E-01 4,66E-01 5,18E-01 0,52E-02 6,29E-01 2,23E-01 -2,41E-01 -2,41E-01 -2,12E-01 -2,01E-01 -1,05E-01	-3.40E-01 1.61E-01 1.06E+00 6.18E-01 1.07E+00 2.25E-01 6.95E-01 -6.95E-01 -5.38E-01 -5.38E-01 -5.38E-01 -7.49E-01 -5.96E-01 1.11E-01 2.77E-01 1.11E-01 2.77E-01 -1.08E+00 -2.32E-01 3.79E-01	2 Unchanged Medium 3 Unchanged Medium 3 Unchanged Medium 4 Unchanged Medium 5 Unchanged Medium 5 Unchanged Medium 6 Unchanged Medium 7 Unchanged Medium 8 Unchanged Medium 9 Unchanged Medium
Q06643 Q95372 Q15046 Q13312 Q09160 P40925 P40925 Q9NQG1 Q9NQG1 Q9NQG1 Q9NQG1 Q9Y5V3 P41223 Q9Y5V3 P43121 Q92494 Q14444 Q9NPE2 Q92571 Q92571 Q92573 Q95204 P80297 Q99735 Q99735 Q99736 Q99736 Q99736 Q99736 Q99736	O21er4 D05cd7 O7er6 J04er7 N06ab6 F09ab7 B17ab7 D17cd3 A05ab6 J23ab7 L01ab7 L01ab7 C15eb7 L09cd4 G13cd6 G13ab7 L09cd4 G13cd6 G13ab7 L09cd4 G13cd6 G13eb7 C15eb7 D12er2 G07ab7 N11er3 H20ab7 C18er7 D13gh8 M14er3 E07er2	Lsm3 protein lung cancer candidate lymphotoxin beta (TNF superfam lysophospholipase II lysosomal-assodated membrane lysyl oxidase-like 1 lysyl-IRNA synthetase MAD1 mitotic arrest deficient-like major histocompatibility complex; malate dehydrogenase 1; NAD (s malate dehydrogenase 2; NAD (s manosyl (alpha-1;3-)-dycoprote manonsyl (alpha-1;3-)-dycoprote maternal G10 transcript melanoma artilgen; family D; 1 melanoma cell adhesion molecul membrane corponent; chromost membrane corponent; chromost mesenchymal stem cell protein C mesoderm specific transcript hon metalloprotease 1 (pitrilysin famil metallothionein 11. mitochondrial darrier homolog 2 mitochondrial ribosomal protein 1 mitochondrial ribosomal protein 2 mitochondrial ribosomal protein 1	1641.224 690.6081 405.7368 480.0562 952.6393 1391.453 900.8177 807.7267 1144.218 1525.837 1253.21 533.2883 619.6733 600.1049 372.2443 397.4495 886.0756 973.362 549.4803 749.6938 617.581 735.157 589.6327 589.6327 589.6328	772 1040 737 2000 1630 556 503 2710 768 1020 1530 800 369 583 952 795 1050 666 932 446 349 502 686 692 1910 421	601.3296 298.9254 752.7079 1277.773 2243.025 539.813 574.4266 1290.783 1089.134 1212.544 1827.301 736.627 887.6535 636.621 575.7695 674.881 432.1718 834.044 64.8174 1085.26 537.173 478.8153 656.4441 173.9779 887.6299 173.2102 17	1511.517 688.0098 582.7217 656.485 1408.92 1753.531 665.5485 628.231 1841.106 1007.157 1251.158 1536.52 689.9075 625.3732 606.4146 503.5179 1008.268 704.4837 952.5469 533.4868 532.52 952.5469 533.4868 532.899 532.747 533.4868 534.4868 534.4868 534.4868 534.4868 534.4868 535.4868 534.4868 53	-3.95E-02 -2.00E-01 -4.41E-01 6.49E-01 4.42E-01 -7.39E-01 -7.12E-02 -3.2E-01 5.18E-01 4.66E-01 5.18E-01 4.66E-01 5.18E-01 -2.23E-01 -2.23E-01 -2.41E-01 -2.23E-01 -2.41E-01 -2.23E-01 -2.41E-01 -2.23E-01 -2.41E-01 -2.41E-01 -2.41E-01 -2.41E-01 -2.41E-01 -1.44E+00 -2.41E-01 -2.4	-3.40E-01 1.61E-01 1.05E-01 1.07E-00 2.25E-01 -6.95E-01 -6.95E-01 -5.30E-01 5.30E-01 5.30E-01 5.85E-01 -7.49E-02 5.96E-01 1.26E+00 -1.56E-01 4.71E-01 -1.00E+00 -2.37E-01 1.49E+00 2.37E-01 1.49E+00 2.37E-01 1.49E+00 2.56E-02	2 Unchanged Medium 3 Unchanged Medium 4 Unchanged Medium 5 Unchanged Medium 5 Unchanged Medium 6 Unchanged Medium 6 Unchanged Medium 7 Unchanged Medium 8 Unchanged Medium 9 Unchanged Medium
Q06643 Q95372 Q15046 Q13312 Q09150 P40925 Q16626 Q9NQG1 P26572 Q10469 P41223 Q9Y5V3 P43121 Q92494 Q9NPE2 Q92571 Q95251 Q99735 Q99735 Q99735 Q997387 Q997387 Q997387 Q997387	O21er4 D05cd7 O7er6 J04er7 ND6ab6 F09ab7 B17ab7 D17cd3 A05ab6 J23ab7 L01ab7 L01ab7 G11ab7 C06gh5 G11ab7 G15ab7 L09cd4 G13cd6 I22er5 D11er1 H15ab7 L09cd4 G13cd6 I22er5 D11er1 H15ab7 C12er2 G07ab7 T11er3 H20ab7 C18er7 M14er3 E07er2 E07er2 E07er8	Lsm3 protein lung cancer candidate lymphotoxin beta (TNF superfam lysophospholipase II lysosomal-associated membrane lysophospholipase II lysosomal-associated membrane lysyl oxidase-like 1 lysyl-IRNA synthetase MAD1 mitotic arrest deficient-like major histocompatibility complex; malate dehydrogenase 1; NAD (s malate dehydrogenase 2; NAD (r mala-enhanced anligen marnosidase; beta A: lysosomal-mannosyl (alpha-1;5-)-glycoprote mannosyl (alpha-1;6-)-glycoprote maternal G10 transcript melanoma antigen; family D; 1 melanoma cell adhesion molecul membrane component; chromosc mesenchymal stem cell protein CD4 membrane component; chromosc mesenchymal stem cell protein Cmesoderm specific transcript hon metalloprotease 1 (plirilysin famil metallothionein 1). microsomal glutathione S-transfemiorobibule-associated protein 1 mitochondrial carrier homolog 2 mitochondrial ribosomal protein 1 m	1641.224 690.6061 405.7368 480.0562 952.6393 1391.453 900.8177 807.7267 1524.076 1524.076 1524.076 1524.076 1619.6733 600.1049 372.2443 372.2443 374.4959 866.0756 973.362 549.4803 749.6938 617.581 735.157 589.6327 527.4681 247.4817 1874.893 401.0368 553.658	772 1040 737 2000 1630 556 503 2710 768 1020 1530 800 369 583 952 795 1050 666 932 446 349 502 686 692 1910 421 421	601.3296 298.9254 752.7079 1277.773 2243.025 539.813 574.4266 1290.783 1089.134 1212.544 1827.301 736.627 887.6535 636.621 575.7695 1674.881 432.1718 834.044 464.8174 1085.26 537.1735 478.8153 656.4441 373.9779 887.6299 1893.062 732.113 887.6299	1511.517 688.0098 582.7217 656.485 1408.92 1753.531 665.5485 628.231 1841.106 1007.157 1251.158 1536.52 689.9075 625.3732 606.4148 503.5179 1008.268 704.4837 952.9144 500.5533 922.569 520.899 582.747 520.9144 782.145 517.9208 521.47 1892.145 517.9208 634.2498	-3.95E-02 -2.00E-01 -4.41E-01 6.49E-01 4.24E-01 6.69E-01 -1.23E-01 -2.40E-01 -7.12E-02 -3.22E-01 5.44E-01 4.66E-01 8.52E-02 6.29E-01 2.29E-01 1.24E-01 5.33E-01 -2.01E-01 -2.5E-01 1.55E-01 1.55E-01 1.55E-01 1.55E-01 1.85E-02 8.68E-01	-3.40E-01 1.61E-01 1.06E+00 6.18EE-01 1.07E+00 6.18EE-01 6.95EE-01 6.95EE-01 6.95EE-01 2.87E-01 5.98EE-01 2.87E-01 5.96E-01 1.16E-01 3.14E-01 4.27E-01 3.14E-01 4.77E-01 3.14E-01 4.79E-02 4.79E-02 4.57E-01	2 Unchanged Medium 3 Unchanged Medium 3 Unchanged Medium 4 Unchanged Medium 5 Unchanged Medium 5 Unchanged Medium 6 Unchanged Medium 7 Unchanged Medium 7 Unchanged Medium 8 Unchanged Medium 9 Unchanged Medium
Q06643 Q95372 Q15046 Q13312 Q09160 P40925 P40926 Q16626 Q9NQG1 P26572 Q10469 P41223 Q9Y5V3 P43121 Q92494 Q14444 Q9NPE2 Q92571 Q95204 Q982W5 Q982W5 Q995W3	O21er4 D05cd7 O7er6 J04er7 N06ab6 N06ab6 N06ab6 N09ab7 N17ab7 D17cd3 A05ab6 J23ab7 L01ab7 L01ab7 L01ab7 L09cd4 G13ab6 L22ef5 D11er1 H15ab7 L09cd4 G13cd6 L22ef5 D11er1 H15ab7 C18er7 D13gh8 M14er3 E07er2 A12er8 M06gh2	Lsm3 protein lung cancer candidate lymphotoxin beta (TNF superfam lysophospholipase II lysosomal-associated membrane lysyl oxidase-like 1 lysyl-IRNA synthetase MAD1 mitotic arrest deficient-like major histocompatibility complex; malate dehydrogenase 1; NAD (s malate dehydrogenase 2; NAD (mala-enhanced antigen mamosidase; beta A: lysosomal-mannosyl (alpha-1;3-)-glycoprote maternal G10 transcript melanoma antigen; family D: 1 melanoma antigen; family D: 1 melanoma collactor protein (CD4 membrane component; chromosc mesenchymal stem cell protein C mesoderm specific transcript hon metalloprotease 1 (pltritysin famil metallothlonein 11. microsomal glutathione S-transfemicrotubule-associated protein 1 milochondrial dearrer hornolog 2 mitochondrial ribosomal protein 1 milochondrial ri	1641.224 690.6061 405.7368 480.0562 952.6393 390.8177 807.7267 1144.218 1525.837 1253.21 533.2883 619.6733 600.1049 372.2443 397.4495 886.0756 973.362 549.48038 617.581 735.157 589.6327 527.4698 247.2817 1874.693 401.0368 553.658 642.9143	772 1040 737 2000 1630 556 503 2710 788 1020 1530 800 369 583 563 952 7955 1050 666 932 446 349 502 686 692 1910 421 760 496	601,3296 298,9254 752,7079 1277,773 2243,025 539,813 574,4266 1290,763 1089,134 1212,544 1827,301 736,627 887,6535 636,621 575,7695 1674,881 432,1718 834,044 464,8174 1085,26 537,1735 478,815 656,4441 173,9779 887,6292 732,113 589,0617 701,1442	1511.517 688.0098 582.7217 656.485 1408.92 1753.531 665.5485 628.231 1841.106 1007.157 1251.158 1536.52 689.9075 625.3732 606.4146 503.5179 1008.268 704.4837 952.9144 560.0583 520.899 522.747 529.1741 609.0447 1892.145 517.9208 634.2498 613.4293	-3,95E-02 -2,00E-01 -4,41E-01 6,49E-01 4,24E-01 -7,39E-01 -4,92E-01 -7,12E-02 -3,32E-01 5,14E-01 4,66E-01 5,14E-01 4,66E-01 5,14E-01 4,66E-01 5,18E-01 0,104E+00 -2,23E-01 -2,41E-01 1,55E-01 -4,96E-01 1,39E-02 0,88E-01 0,88E-01 0,88E-01 0,88E-01 1,39E-02 1,25E-01	-3.40E-01 1.61E-01 1.06E+00 6.18E-01 1.07E+00 2.25E-01 6.95E-01 -6.95E-01 -5.38E-01 -5.38E-01 -5.38E-01 -5.38E-01 -7.49E-01 5.56E-01 1.11E-01 2.76E-01 1.11E-01 2.71E-01 -1.08E+00 2.56E-02 6.88E-02 6.88E-02 6.88E-03 -7.49E-01	2 Unchanged Medium 3 Unchanged Medium 3 Unchanged Medium 3 Unchanged Medium 4 Unchanged Medium 5 Unchanged Medium 5 Unchanged Medium 6 Unchanged Medium 7 Unchanged Medium 8 Unchanged Medium 9 Unchanged Medium
Q06643 Q95372 Q15046 Q13312 Q09160 P40925 P40925 Q9NQG1 Q9NQG1 P26572 Q10469 P41223 Q9Y5V3 P43121 Q92494 Q14444 Q3NPE2 Q92571 Q92571 Q92573 Q99750 Q9	O21er4 D05cd7 O7cef6 J04er7 N06ab6 F09ab7 B17ab7 D17cd3 A05ab6 J23ab7 L01ab7 L01ab7 L05eb7 L0	Lsm3 protein lung cancer candidate lymphotoxin beta (TNF superfam lysophospholipase II lysosomal-assodated membrane lysyl oxidase-like 1 lysyl-IRNA synthetase MAD1 mitotic arrest deficient-like major histocompatibility complex, malate dehydrogenase 1; NAD (s malate dehydrogenase 2; NAD (s malate dehydrogenase 1; NAD (s malate dehydrogenase 1; NAD (s malate dehydrogenase 2; NAD (s malate dehydrogenase 2; NAD (s malate dehydrogenase 2; NAD (s malate dehydrogenase 1; NAD (s manosyl (alpha-1;3-)-dycoprote manonsyl (alpha-1;3-)-dycoprote malernal G10 transcript melanoma artilgen; family D; 1 melanoma cell adhesion molecul membrane corponent; chromost mesenchymal stem cell protein C mesoderm specific transcript hon metalloprotease 1 (pitrilysin famil metallothoneria 1; mitochondrial dibosomal protein 1 mitochondrial ifbosomal protein 1 mitochondrial ribosomal protein 1	1641.224 690.6081 405.7368 480.0562 952.6393 1391.453 900.8177 807.7267 1144.218 1525.837 1253.21 533.2883 619.6733 600.1049 372.2443 397.4495 886.0756 973.362 549.4803 617.581 735.157 589.6327 527.4698 247.2817 1874.893 401.0368 553.658 642.9143 741.5383	772 1040 737 2000 1630 556 503 2710 768 1020 1530 800 369 583 952 795 1050 666 932 446 349 502 686 692 1910 421 760 496	601.3296 298.9254 752.7079 1277.773 2243.025 539.813 574.4266 1290.783 1089.134 1212.544 1827.301 736.627 87.6535 636.621 575.7695 1674.881 432.1718 834.044 464.8174 1085.26 537.1735 478.8153 656.4441 373.9779 887.6299 1893.062 737.9779 887.6299 1893.0617 701.1442 826.8816	1511.517 688.0098 582.7217 656.485 1408.92 1753.531 665.5485 628.231 1841.106 1007.157 1251.158 1536.52 689.9075 625.3732 606.4146 503.5179 1008.268 704.4837 922.5469 533.4868 520.899 522.5469 533.4868 520.899 522.5469 533.4868 536.357 53	-3.95E-02 -2.00E-01 -4.41E-01 6.49E-01 4.42E-01 -7.39E-01 -7.12E-02 -3.2E-01 5.18E-01 6.69E-01 5.18E-01 4.66E-01 5.18E-01 6.29E-01 2.01E-01 -2.21E-01 -2.21E-01 1.39E-02 1.39E-02 1.29E-01	-3.40E-01 1.61E-01 1.65E-00 6.18E-01 1.07E+00 2.25E-01 -6.96E-01 -6.96E-01 1.5.36E-01 2.07E-01 2.07E-01 2.07E-01 2.07E-01 1.26E+00 -1.56E-01 1.11E-01 -1.11E-01 -1.08E+00 -2.32E-01 1.49E+00 2.56E-02 4.57E-01 -3.74E-01 -3.74E-01	2 Unchanged Medium 3 Unchanged Medium 4 Unchanged Medium 5 Unchanged Medium 5 Unchanged Medium 6 Unchanged Medium 7 Unchanged Medium 7 Unchanged Medium 8 Unchanged Medium 9 Unchanged Medium
Q06643 Q95372 Q15046 Q13312 Q09150 P40925 Q16626 Q16626 Q16626 Q950G1 Q95733 P41223 Q95741 Q92494 Q9571 Q9574 Q99735	021er4 D05cd7 D07er6 J04er7 ND6ab6 F09ab7 B17ab7 D17cd3 A05ab6 J23ab7 L01ab7 G11ab7 G15ab7 L09cd4 G13cd6 I23er5 D11er1 H15ab7 L09cd4 G13cd6 I22er5 D11er1 H15ab7 C12er2 G07ab7 H15ab7 L09cd4 G13cd6 I22er5 D11er1 H15ab7 L09cd4 G13cd6 I22er5 D11er1 H15ab7 L09cd4 G13cd6 I22er2 G07ab7 H16ab7	Lsm3 protein lung cancer candidate lymphotoxin beta (TNF superfam lysophospholipase II lysosomal-associated membrane lysophospholipase II lysosomal-associated membrane lysyl oxidase-like 1 lysyl-IRNA synthetase MAD1 mitotic arrest deflicient-like major histocompatibility complex; malate dehydrogenase 1; NAD (s malate dehydrogenase 2; NAD (r mala-enhanced antigen mamnosidase; beta A: lysosomal- mannosyl (alpha-1;5-)-glycoprote mannosyl (alpha-1;6-)-glycoprote maternal G10 transcript melanoma antigen; family D: 1 melanoma cell adhesion molecut membrane component; chromosc mesenchymal stem cell protein (CD4 membrane component; chromosc mesenchymal stem cell protein E mesoderm specific transcript hon metalloprotease 1 (plirilysin famil metallothionein 11. microsomal glutathione S-transfe mirothodrial enter homolog 2 mitochondrial ribosomal protein 1	1641.224 690.6061 405.7368 480.0562 952.6393 1391.453 900.8177 807.7267 1544.218 1525.837 1253.21 533.2833 610.1049 372.2443 397.4495 886.0756 973.362 549.4803 749.6938 617.581 735.157 589.6327	772 1040 737 2000 1630 556 503 2710 768 1020 1530 800 369 583 952 795 1050 666 932 446 349 502 686 692 1910 421 760 496 1000 910	601.3296 298.9254 752.7079 1277.773 2243.025 539.813 574.4266 1290.783 1089.134 1212.544 1827.301 736.627 887.6535 636.621 575.7695 1674.881 432.1718 834.044 464.8174 1085.26 537.1735 478.8153 656.4441 373.9779 887.6299 1893.062 732.113 589.0617 701.1442 826.8816 926.8816 926.8816	1511.517 688.0098 582.7217 656.485 1408.92 1753.531 665.5485 628.231 1841.106 1007.157 1251.158 1536.52 689.9075 625.3732 606.4146 503.5179 1008.268 704.4837 952.9144 500.5533 922.5469 533.4660 520.899 582.747 1599.7441 1699.7	-3.95E-02 -2.00E-01 -4.41E-01 -4.41E-01 -4.92E-01 -7.39E-01 -4.92E-01 -7.12E-02 -3.32E-01 -3.32E-01 -5.18E-01 -8.52E-02 -2.29E-01 -2.29E-01 -2.32E-01 -2.32E-01 -1.04E+00 -1.04E+00 -1.04E+00 -1.04E-01 -2.52E-01 -1.55E-01 -1.86E-01 -1.86E-01 -1.57E-01 -1.57E-01 -1.57E-01	-3.40E-01 1.61E-01 1.06E+00 6.18EE-01 1.07E+00 6.98EE-01 6.98EE-01 6.98EE-01 2.87E-01 5.38EE-01 2.87E-01 5.38EE-01 1.58EE-01 1.58EE-01 1.58EE-01 1.56E-01 3.14E-01	2 Unchanged Medium 3 Unchanged Medium 3 Unchanged Medium 3 Unchanged Medium 4 Unchanged Medium 4 Unchanged Medium 5 Unchanged Medium 6 Unchanged Medium 6 Unchanged Medium 7 Unchanged Medium 8 Unchanged Medium 9 Unchanged Medium
Q06643 Q95372 Q15046 Q13312 Q09160 P40925 P40926 Q16626 Q10469 P41223 Q975V3 P43121 Q92494 Q14444 Q9NPE2 Q92571 Q95204 Q95095 Q9	O21er4 D05cd7 O7cr6 J04er7 N06ab6 N06ab6 N06ab6 N09ab7 N17ab7 D17cd3 A05ab6 J23ab7 L01ab7 L01ab7 L01ab7 L01ab7 L02ed4 G13ab6 G13ab7 L09cd4 G13cd6 H22ef5 D11er1 H15ab7 C18er7 D13gh8 M14er3 E07er2 A12er8 M06gh2 O13er8 O04er2 CK18er2	Lsm3 protein lung cancer candidate lymphotoxin beta (TNF superfam lysophospholipase II lysosomal-associated membrane lysyl oxidase-like 1 lysyl-IRNA synthetase MAD1 mitotic arrest deficient-like major histocompatibility complex; malate dehydrogenase 1; NAD (s malate dehydrogenase 2; NAD (mala-enhanced antigen mamosidase; beta A: lysosomal- mannosyl (alpha-1;3-)-glycoprote maternal G10 transcript melanoma antigen; family D; 1 melanoma antigen; family D; 1 melanoma collactor protein (CD4 membrane component; chromost mesenchymal stem cell protein C mesoderm specific transcript hon metalloprotease 1 (pltritysin famil metallothlonein 11. microsomal glutathione S-transfe microtubule-associated protein 1 mitochondrial ribosomal protein 1	1641.224 690.6061 405.7368 480.0562 952.6393 390.8177 807.7267 1144.218 1525.837 1253.21 533.2883 619.6733 600.1049 372.2443 397.4495 886.0756 973.362 549.4803 617.581 735.157 589.6327 527.4698 247.2817 1874.893 401.0368 553.658 642.9143 741.5383 914.9722 1217.101	772 1040 737 2000 1630 556 503 2710 788 1020 1530 800 369 583 563 952 7955 1050 666 932 446 349 502 686 692 1910 421 760 496 1000 910 891	601,3296 298,9254 752,7079 1277,773 2243,025 539,813 574,4266 1290,763 1089,134 1212,544 1827,301 736,627 887,6535 636,621 575,7695 1674,881 432,1718 834,044 464,8174 1085,26 537,1735 478,815 656,4441 737,9779 887,6299 1893,0617 701,1442 286,8816 956,7888 1196,143	1511.517 688.0098 582.7217 656.485 1408.92 1753.531 665.5485 628.231 1841.106 1007.157 1251.158 1536.52 689.9075 625.3732 606.4146 503.5179 1008.268 704.4837 952.9144 500.5689 532.868 520.899 582.747 599.1741 609.0447 1892.145 517.208 634.2498 634.2498 634.2498 634.2498 634.2498 634.2498 634.2498 634.2498 634.2498 636.3674 927.2731 1101.306	-3,95E-02 -2,00E-01 -4,41E-01 6,49E-01 4,24E-01 -7,39E-01 -4,92E-01 -7,12E-02 -3,32E-01 5,44E-01 4,66E-01 5,18E-01 4,66E-01 5,18E-01 2,08E+00 -2,21E-01 -2,01E-01 -6,19E-01 1,04E+00 1,25E-02 0,68E-01 1,04E+00 1,04E-01 1,	-3.40E-01 1.61E-01 1.06E+00 6.18E-01 1.07E+00 2.25E-01 6.95E-01 -6.95E-01 -5.38E-01 -5.38E-01 -5.38E-01 -5.38E-01 1.56E-01 1.56E-01 1.26E-01 1.11E-01 2.77E-01 3.74E-01 -1.47E-01 -1.47E-01 -1.47E-01 -1.47E-01 -1.47E-01 -1.47E-01 -1.47E-01 -1.77E-01 4.77E-01 4.77E-01 4.77E-01	2 Unchanged Medium 3 Unchanged Medium 3 Unchanged Medium 3 Unchanged Medium 4 Unchanged Medium 5 Unchanged Medium 5 Unchanged Medium 5 Unchanged Medium 6 Unchanged Medium 7 Unchanged Medium 8 Unchanged Medium 8 Unchanged Medium 9 Unchanged Medium
Q06643 Q95372 Q15046 Q13312 Q09160 P40925 P40926 Q1626 Q1626 Q9NQG1 P26572 Q10469 P41223 Q9Y5V3 P43121 Q92494 Q14444 Q3NPE2 Q92571 Q95204 P80297 Q99735 Q99735 Q99735 Q99736 Q99736 Q99736 Q99736 Q99737	O21er4 D05cd7 O7cef6 J04er7 N06ab6 F09ab7 B17ab7 D17cd3 A05ab6 J23ab7 L01ab7 L01ab7 L01ab7 L01ab7 L05cd4 G13cd6 G13cd6 G13cd6 G13cd7 L01ab7 C16er7 D13gh8 M14er3 E07er2 A12er8 M06gh2 O13er8 O04er2 K18er2 M15er2 M15er2	Lsm3 protein lung cancer candidate lymphotoxin beta (TNF superfam lysophospholipase II lysosomal-associated membrane lysyl oxidase-like 4 lysyl-IRNA synthetase MAD1 mitotic arrest deficient-like rnajor histocompatibility complex; malate dehydrogenase 1; NAD (s malate dehydrogenase 2; NAD (s malate dehydrogena	1641.224 690.6061 405.7368 480.0562 952.6393 1391.453 900.8177 807.7267 1524.076 1144.218 1525.837 1253.21 533.2883 619.6733 600.1049 372.2443 397.4495 886.0756 973.362 549.4803 749.6938 617.581 735.157 589.6327 527.4698 247.2817 1874.893 401.0368 553.658 642.9143 741.5393 314.9722 1217.101	772 1040 737 2000 1630 556 503 2710 768 1020 1530 800 369 583 952 795 1050 666 932 446 349 502 1910 686 692 1910 496 1000 910 910	601.3296 298.9254 752.7079 1277.773 2243.025 539.813 574.4266 1290.783 1089.134 1212.544 1827.301 736.627 887.6535 636.621 575.7695 1674.881 432.1718 834.044 464.8174 1085.26 537.1735 478.8153 656.4441 373.9779 887.6299 1893.062 737.9779 887.6299 1893.0617 701.1442 696.7888 1196.143 1372.352	1511.517 688.0098 582.7217 656.485 1408.92 1753.531 665.5485 628.231 1841.106 1007.157 1251.158 1536.52 689.9075 625.3732 606.4146 503.5179 1008.268 704.4837 952.5469 533.4868 520.899 582.747 1591.741 690.447 1892.145 517.9208 634.2498 613.4293 6	-3.95E-02 -2.00E-01 -4.41E-01 6.49E-01 4.42E-01 -7.39E-01 -7.39E-01 -7.12E-02 -3.24E-01 5.18E-01 4.66E-01 5.18E-01 4.66E-01 5.18E-01 -2.23E-01 -2.23E-01 -2.41E-01 -2.23E-01 -2.41E-01 -2.23E-01 -2.41E-01 -2.01E-01 -2.51E-01 1.39E-02 1.25E-01 6.45E-02 -2.51E-01 6.52E-02 -2.51E-01 -2.51E-01 6.52E-02 -2.51E-01 -2.51E-01	-3.40E-01 1.61E-01 1.65E-00 6.18E-01 1.07E+00 2.25E-01 -6.95E-01 -6.95E-01 -5.38E-01 2.07E-01 2.07E-01 2.07E-01 2.07E-01 1.26E+00 -1.56E-01 1.11E-01 -1.11E-01 -1.08E+00 -2.32E-01 1.49E+00 2.56E-02 4.57E-01 -3.74E-01 -1.77E-03 -3.74E-01 -1.77E-03 -3.74E-01	2 Unchanged Medium 3 Unchanged Medium 3 Unchanged Medium 4 Unchanged Medium 5 Unchanged Medium 5 Unchanged Medium 5 Unchanged Medium 5 Unchanged Medium 6 Unchanged Medium 7 Unchanged Medium 8 Unchanged Medium 8 Unchanged Medium 9 Unchanged Medium
Q06643 Q95372 Q15046 Q13312 Q09160 P40925 P40926 Q16626 Q10469 P41223 Q975V3 P43121 Q92494 Q14444 Q9NPE2 Q92571 Q95204 Q95095 Q9	O21er4 D05cd7 O7cr6 J04er7 N06ab6 N06ab6 N06ab6 N09ab7 N17ab7 D17cd3 A05ab6 J23ab7 L01ab7 L01ab7 L01ab7 L01ab7 L02ed4 G13ab6 G13ab7 L09cd4 G13cd6 H22ef5 D11er1 H15ab7 C18er7 D13gh8 M14er3 E07er2 A12er8 M06gh2 O13er8 O04er2 CK18er2	Lsm3 protein lung cancer candidate lymphotoxin beta (TNF superfam lysophospholipase II lysosomal-associated membrane lysyl oxidase-like 1 lysyl-IRNA synthetase MAD1 mitotic arrest deficient-like major histocompatibility complex; malate dehydrogenase 1; NAD (s malate dehydrogenase 2; NAD (mala-enhanced antigen mamosidase; beta A: lysosomal- mannosyl (alpha-1;3-)-glycoprote maternal G10 transcript melanoma antigen; family D; 1 melanoma antigen; family D; 1 melanoma collactor protein (CD4 membrane component; chromost mesenchymal stem cell protein C mesoderm specific transcript hon metalloprotease 1 (pltritysin famil metallothlonein 11. microsomal glutathione S-transfe microtubule-associated protein 1 mitochondrial ribosomal protein 1	1641.224 690.6061 405.7368 480.0562 952.6393 1391.453 900.8177 807.7267 1524.076 1144.218 1525.837 1253.21 533.2883 619.6733 600.1049 372.2443 397.4495 886.0756 973.362 549.4803 749.6938 617.581 735.157 589.6327 527.4698 247.2817 1874.893 401.0368 553.658 642.9143 741.5393 314.9722 1217.101	772 1040 737 2000 1630 556 503 2710 768 1020 1530 800 369 583 952 795 1050 666 932 446 349 502 1910 686 692 1910 496 1000 910 910	601.3296 298.9254 752.7079 1277.773 2243.025 539.813 574.4266 1290.783 1089.134 1212.544 1827.301 736.627 887.6535 636.621 575.7695 1674.881 432.1718 834.044 464.8174 1085.26 537.1735 478.8153 656.4441 373.9779 887.6299 1893.062 737.9779 887.6299 1893.0617 701.1442 696.7888 1196.143 1372.352	1511.517 688.0098 582.7217 656.485 1408.92 1753.531 665.5485 628.231 1841.106 1007.157 1251.158 1536.52 689.9075 625.3732 606.4146 503.5179 1008.268 704.4837 952.5469 533.4868 520.899 582.747 1591.741 690.447 1892.145 517.9208 634.2498 613.4293 6	-3.95E-02 -2.00E-01 -4.41E-01 6.49E-01 4.42E-01 -7.39E-01 -7.39E-01 -7.12E-02 -3.24E-01 5.18E-01 4.66E-01 5.18E-01 4.66E-01 5.18E-01 -2.23E-01 -2.23E-01 -2.41E-01 -2.23E-01 -2.41E-01 -2.23E-01 -2.41E-01 -2.01E-01 -2.51E-01 1.39E-02 1.25E-01 6.45E-02 -2.51E-01 6.52E-02 -2.51E-01 -2.51E-01 6.52E-02 -2.51E-01 -2.51E-01	-3.40E-01 1.61E-01 1.06E+00 6.18E-01 1.07E+00 2.25E-01 6.95E-01 -6.95E-01 -5.38E-01 -5.38E-01 -5.38E-01 -5.38E-01 1.56E-01 1.56E-01 1.26E-01 1.11E-01 2.77E-01 3.74E-01 -1.47E-01 -1.47E-01 -1.47E-01 -1.47E-01 -1.47E-01 -1.47E-01 -1.47E-01 -1.77E-01 4.77E-01 4.77E-01 4.77E-01	2 Unchanged Medium 3 Unchanged Medium 3 Unchanged Medium 3 Unchanged Medium 4 Unchanged Medium 5 Unchanged Medium 5 Unchanged Medium 5 Unchanged Medium 6 Unchanged Medium 7 Unchanged Medium 8 Unchanged Medium 8 Unchanged Medium 9 Unchanged Medium

Q9BYC		mitochondrial ribosomal protein	1 490.5463	665	619,0928	591,611	8 3.36E-01	4.39E-01	2 Unchanged Medium
Q9Y3D3 Q9Y676		milochondrial ribosomal protein	5 966.7611	1020					2 Unchanged Medium
Q9Y3D5		milochondrial ribosomal protein milochondrial ribosomal protein		580 1080					2 Unchanged Medium
P82921		mitochondrial ribosomal protein		1190			1 2.46E-01 3 5.76E-01		2 Unchanged Medium
P82650		milochondrial ribosomal protein		515			1 -2.36E-01		2 Unchanged Medium
Q9Y2Q9		milochondrial ribosomal protein	1439.076	1420			-1.14E-01		2 Unchanged Medium 2 Unchanged Medium
Q9NP92		mitochondrial ribosomal protein	5 669.1652	532			5 -1.16E-01		2 Unchanged Medium
Q02750 Q02779		milogen-activated protein kinase	615,689	701		590,1129	9 -4.40E-D1	1.87E-01	2 Unchanged Medium
Q12851		mitogen-activated protein kinase mitogen-activated protein kinase	988.0971	423			-2.29E-01		2 Unchanged Medium
P49137	J06ef5	mitogen-activated protein kinase		1020 1000			-1.67E-02		2 Unchanged Medium
Q9BYG3		MKI67 (FHA domain) Interacting		407			1.81E-01 2 -1.36E+00		2 Unchanged Medium
Q15014		MORF-related gene X	1146.48	1030				-1.61E-01	2 Unchanged Medium
Q15773	F22ef4	mRNA; clane:PO2ST9	1052.442	1040	905.2614		-2.17E-01	-1.60E-02	2 Unchanged Medium 2 Unchanged Medium
P35579	M18ef5 001ef1	myeloid leukernia factor 2	577.4585	467				-3.06E-01	2 Unchanged Medium
Q9Y6D2		myosin; heavy polypeptide 9; no N-acelyltransferase 5 (ARD1 hor	1426,003	1150			-2.44E-01		2 Unchanged Medium
P41227	G13gh1	N-acetyltransferase; homolog of	800.6676	708 905					2 Unchanged Medium
Q13510	H20ab2	N-acylsphingosine amldohydrola		1660				1.76E-01 1.14E+00	2 Unchanged Medium
043678	O09ab7	NADH dehydrogenase (ubiquino	1366.754	1930					2 Unchanged Medium 2 Unchanged Medium
O95167 P56556	E24ab7	NADH dehydrogenase (ubiquino	1431.655	1770					2 Unchanged Medium
O95182	O11ab7 K06ab7	NADH dehydrogenase (ubiquino		916			-1.05E+00	-8.49E-01	2 Unchanged Medium
P51970	L04cd8	NADH dehydrogenase (ubiquino NADH dehydrogenase (ubiquino		777	725.1711		-4.87E-02		2 Unchanged Medium
O96000	J10ab8	NADH dehydrogenase (ubiquino	1110 197	728 1440	1021.163 1111.615				2 Unchanged Medium
O95178	J06ab8	NADH dehydrogenase (ubiquino	925,3926	768	815.266			3.71E-01	2 Unchanged Medium
095139	M03ab8	NADH dehydrogenase (ubiquino	1392,873	1480	2162,279		6.34E-01		2 Unchanged Medium 2 Unchanged Medium
P17568	H02ab8	NADH dehydrogenase (ubiquino	787.0681	567	591.1924	648.3703	-4.13E-01		2 Unchanged Medium
Q9Y6M9 O43677	P16ab8 M05ab8	NADH dehydrogenase (ubiquino	1299,803	1450	746,8017	1164,524	-7.99E-01	1.55E-01	2 Unchanged Medium
000217	M09ab8	NADH dehydrogenase (ubiquino NADH dehydrogenase (ubiquino	1392 411	601	532,551		-1.39E-01		2 Unchanged Medium
P19404	M07ef1	NADH dehydrogenase (ubiquino	1151.224	1130 443	1194,001 356,308	650 0045	-2.12E-01 -1.69E+00	-2.96E-01	2 Unchanged Medium
076008	A12cd6	Nef-associated factor 1	720,2744	951	424.04	698,4474		4.01E-01	2 Unchanged Medium
Q9UPY4		nesca protein	425,9149	469	621,9336	505,6782		1.40E-01	2 Unchanged Medium 2 Unchanged Medium
Q99742	L19gh8 007ab8	neurabin il	1241.544	862	1023,077	1042.081	-2.79E-01	-5.27E-01	2 Unchanged Medium
060448	P20ef3	neuronal PAS domain protein 1 neuronal thread protein	1076.259	405	606.3589				2 Unchanged Medium
Q9UNW9		neuro-oncological ventral antiger	1831,592 1049,959	1660 1050	1490.624 1219,792	1662.051	-2.97E-01		2 Unchanged Medium
Q9UGL9	C09gh3	NICE-1 protein	1026.838	1360	381,058		2.16E-01 -1.43E+00	1.70E-03 4.08E-01	2 Unchanged Medium
Q9UGL6	M09ef4	NICE-5 protein	912.7542	1390	955.016	1086,502		6.09E-01	2 Unchanged Medium 2 Unchanged Medium
Q92982 Q9BPW8	109ef7	ninjurin 1	458,23	689	412.0181	519,6952		5.88E-01	2 Unchanged Medium
Q9NQR4		nipsnap homolog 1 (C. elegans) Nit protein 2	829,4266	529	562,9689	640,3075	-5.59E-01	-6.50E-01	2 Unchanged Medium
Q9UF10	L08ef2	non-canonical ubquitin conjugatis	519,1587 1336,406	868 2000	665,8632	684,4701	3.59E-01	7.42E-01	2 Unchanged Medium
Q9Y385	H11ef2	non-canonical ubquitin conjugati	625.6356	808	1488.008 669.812	1608.811 701.1992	1.55E-01 9.84E-02	5.83E-01	2 Unchanged Medium
Q15233	O02ef1	non-POU domain containing; oct.	1083,363	1210	1234,216	1174,503		3.69E-01 1.55E-01	2 Unchanged Medium 2 Unchanged Medium
Q13137 Q16236	A03cd6 M03cd1	nuclear domain 10 protein	910.7725	914	944.6079	923.1712		5.31E-03	2 Unchanged Medium 2 Unchanged Medium
P08651	B07ef1	nuclear factor (erythroid-derived	1233.751	1180	1043,638	1153.088	-2.41E-01	-6.20E-02	2 Unchanged Medium
O95134	P10cd3	nuclear factor I/C (CCAAT-bindin nuclear localization signal delete	465 2033	624 522	659.6305		-1,38E-01		2 Unchanged Medium
Q15772	N19ab2	nuclear protein; marker for differe	1473.704		536.7333 1157.202	508,0938 1184,738	2.06E-01 -3.49E-01	1.67E-01 -6.75E-01	2 Unchanged Medium
Q9UHY1	L15cd8	nuclear receptor binding protein	515.949	948	619.7512	694,4539	2.64E-01	8.77E-01	2 Unchanged Medium 2 Unchanged Medium
Q9UPC9 Q15325	H23ab2	nuclear receptor coactivator 3	822,7768	1190	932.9058	982.77	1.81E-01	5.36E-01	2 Unchanged Medium
O00567	N22cd2 D05cd6	nuclease sensitive element bindi nucleolar protein 5A (56kD with)	1644.614	1720	2064.24	1808.771	3.28E-01	6.25E-02	2 Unchanged Medium
Q9NX24	A10gh3		659.4335 466.2539		623.2536		-8.14E-02		2 Unchanged Medium
P06748	K06gh6		1605.368	1080	594.8443 1136.972	561,9547 1275,358	3.51E-01 -4.98E-01	4.22E-01	2 Unchanged Medium
P50583	B01ab2	nudix (nucleoside diphosphate lis	1590,763		251,3768		-2.66E+00		2 Unchanged Medium 2 Unchanged Medium
Q9UHM6 Q9Y218	N20gh8	opsin 4 (melanopsin)	1076.73		1643,887	1305.239	6.10E-01	1.50E-01	2 Unchanged Medium 2 Unchanged Medium
Q16612	P22gh1 001cd5	optineurin P311 protein	1014.697		912,1917		-1.54E-01	9.66E-01	2 Unchanged Medium
P55771	J21cd7	paired box gene 9	322.7444 692.508	349 380	2231.846			1.12E-01	2 Unchanged Medium
Q9UKJ1	N21cd8	paired Immunoglobultn-like recer			573.135 558.3225		-2.73E-01 -2.72E-01		2 Unchanged Medium
Q15165	P24ab7	paraoxonase 2	338,5244				7.86E-01		2 Unchanged Medium 2 Unchanged Medium
Q07002	P06ef5	PCTAIRE protein kinase 3	836,5659			795.1858		1.31E-01	2 Unchanged Medium 2 Unchanged Medium
Q9Y3C6 P30405	J07ef2 F14cd5	peptidylprolyl isomerase (cyclop)	548,5938			602.0621		4.14E-01	2 Unchanged Medium
O43924	G24ab8	peplidylprolyl isomerase F (cyclo phosphodiesterase 6D; cGMP-sp	272 +209				-8.50E-01	1.33E-01	2 Unchanged Medium
P17858	K10ab8		950.5746	630 525		567,4949	9.06E-01 -2.95E-01	7.56E-01	2 Unchanged Medium
Q01813	C17ef1	-k- 1 a 111 111.	573,5211			591,6879		1.71E-01	2 Unchanged Medium
000511	N20cd3	phosphoprolein enriched in astro	1330,005					1.70E-01	2 Unchanged Medium 2 Unchanged Medium
P11216 P00749	B16ab8	phosphorylase; glycogen; brain	516.7884	495	988,163		9.35E-01 -		2 Unchanged Medium
P13797	P19ef7 C13cd1	plasminogen activator, urokinase					-5.98E-02		2 Unchanged Medium
Q15102	E16ab8	plastin 3 (T isoform) platelet-activating factor acetylhy	990,1173 469 9778			1062.438	4.93E-01 ·		2 Unchanged Medium
P01127	A08ef5	platelet-derived growth factor bel	500.6466		468.9765 458.4576	537.6672 543.5869		5.20E-01	2 Unchanged Medium
Q15795	C19ef6	pleckstrin homology; Sec7 and o	578.295		518.8327			4.28E-01 3.55E-01	2 Unchanged Medium
043660	O18ab8	pleiotropic regulator 1 (PRL1horr	743.7263		804,3456		1.13E-01		2 Unchanged Medium 2 Unchanged Medium
Q15155 Q9H361	G12ef3	pM5 protein	664,2511	423	490.295	525.8869	-4.38E-01 -	6.51E-01	2 Unchanged Medium 2 Unchanged Medium
	L04gh7 A11cd1	poly(A) binding prolein; cytoplasi poly(rC) binding prolein 2		491	710.9543	653,8286	-9.52E-02 -	6.29E-01	2 Unchanged Medium
		polymerase (RNA) il (DNA direct	1951.241 552 3545		2128.585		1.26E-01 -		2 Unchanged Medium
P52434	F05gh6	polymerase (RNA) II (DNA direct	949.7064		698,4965 1210.411	593.4201 1211.761	3.39E-01 - 3.50E-01		2 Unchanged Medium
	A20cd1	polymerase (RNA) II (DNA direct	1603.557			1613.004		1.84E-01	2 Unchanged Medium
P53803	C17cd1	polymerase (RNA) II (DNA direct	437.9544				2.70E-01		2 Unchanged Medium 2 Unchanged Medium
									- o

Q9UQQ3	J08cd6	POP4 (processing of precursor;	480,5848	645	551,3661	558,9966	1.98E-01	4.25E-01	2 Unchanged Medium
Q13670	B07ab8	postmelotic segregation increase		779	532,0548		-6.11E-01		2 Unchanged Medium
Q9P0J7	L07gh4	potassium channel modulatory fa	1401,591	1720	1192,204		-2.33E-01	2.97E-01	2 Unchanged Medium
060925	K02ab8	prefoldin 1	1631,996	1720	1860,632	1737.289	1.89E-01	7.51E-02	2 Unchanged Medium
Q9UHV9	J15ef4	prefoldin 2	514,4897	882		743,3158	6.95E-01	7.78E-01	2 Unchanged Medium
Q13519	J07ef6	prepronociceplin	564.2079	464	887,4522	638,4071		-2.83E-01	2 Unchanged Medium
P04156	A13ab8	prion protein (p27-30) (Creutzfeli		631	701,8217	710.1438	-1.85E-01		2 Unchanged Medium
Q9UI73	K03ef8	PRO0246 protein	430,412	535	548,8599	504.0018	3.45E-01	3.13E-01	2 Unchanged Medium
Q9P0T3	D14ef1	proapoptotic caspase adaptor pri		1650	2553.652	1820.221	1.02E+00		2 Unchanged Medium
Q02809	P20ab7	procollagen-lysine; 2-oxoglutarat	503,7557	477	1993.232	991.2271	1.98E+00		2 Unchanged Medium
P07237	G13ab8	procollagen-proline; 2-oxoglutars	633,6309	406	575.6635		-1.39E-01		2 Unchanged Medium
P09466	E12ab8	progestagen-associated endome		625	708.8717				2 Unchanged Medium
Q16342	G22ef8	programmed cell death 2	735.0961	759	490,3938		-5.84E-01	4.52E-02	2 Unchanged Medium
O75340	102cd8	programmed cell death 6	1055.886	1090	1356.597	1167,782	3.62E-01	4.70E-02	2 Unchanged Medium
P35232	C22ef5	prohibitin	2045.693	1810	1600.236		-3.54E-01		2 Unchanged Medium
Q9UQ80	003cd1	proliferation-associated 2G4; 389	1902.387	1390	2003,135	1763.59		-4.58E-01	2 Unchanged Medium
Q12796	E05ab3	proline rich 2	1927.78	1530	1475.973		-3.85E-01		2 Unchanged Medium
Q15188	P20cd2	proline-rich protein BstNI subfam	997,5582	1290	1404.4	1232.034	4.93E-01	3.76E-01	2 Unchanged Medium
P3599B	L09ab8	proteasome (prosome; macropaii		1200	1059,594	1090.401	6.85E-02	2.49E-01	2 Unchanged Medium
P47210	N16cd2	proteasome (prosome; macropali	1037.339	1150	1193	1127,919	2.02E-01	1.53E-01	2 Unchanged Medium
O00495	N03ab8	proteasome (prosome; macropali	1043,875	498	563.7775	701,7628	-8.89E-01	-1.07E+00	2 Unchanged Medium
075831	N07ab8	proteasome (prosome; macropali		622	506.8297	630,7237	-5.92E-01	-2.97E-01	2 Unchanged Medium
Q13200	L15ab8	proteasome (prosome; macropali		594	331,4288	523,3702	-9.59E-01	-1.16E-01	2 Unchanged Medium
043242	L17ab8	proteasome (prosome; macropaii		1230	1637.67	1675,386	-3.99E-01	-8.13E-01	2 Unchanged Medium
P55036	N18cd2	proleasome (prosome; macropali		724	588,5346	591.0419	3.53E-01	6.51E-01	2 Unchanged Medium
P48556	L21ab8	proteasome (prosome; macropali		1030	1685,116	1317.42	4.42E-01	-2.72E-01	2 Unchanged Medium
Q92530	G07cd6	proteasome (prosome; macropali		1570	1800.731	1724,253	-3.24E-03	-2.04E-01	2 Unchanged Medium
P28066	J13ab8	proteasome (prosome; macropali		767	775.2704	769,7417	1.62E-02	1.46E-03	2 Unchanged Medium
014818	J15ab8	proteasome (prosome; macropali		1540	1139.994		-1.24E-01	3.12E-01	2 Unchanged Medium
P49721	J19ab8	proteasome (prosome; macropali		2110	1617,062	1853.89	-1.81E-01	2.04E-01	2 Unchanged Medium
P28062	H18ab8	proteasome (prosome; macropaii		905	505.3638			1.82E-01	2 Unchanged Medium
P28065 P10619	B06ef7	proteasome (prosome; macropali		1630	1002.72	1133.626		1.08E+00	2 Unchanged Medium
Q9UNN8	A07ab8	protective protein for beta-galact		907	980,3046	1142.279	-8.51E-01		2 Unchanged Medium
Q15084	F03cd6	protein C receptor; endothelial (E		702	488,5084		-8.69E-01		2 Unchanged Medium
Q9Y2B9	H02cd5 C22cd7	protein disulfide Isomerase-relate		407	859.0228	782.4339	-3.32E-01		2 Unchanged Medium
Q13517	A09ef5	protein kinase (cAMP-dependent		1040	754.6101	814.4047	2.13E-01	6.73E-01	2 Unchanged Medium
P54619	F15ab8	protein kinase C binding protein protein kinase; AMP-activated; g:		587	630,5497	532,9442	7.27E-01	6.25E-01	2 Unchanged Medium
P17612	F13ab8	protein kinase; cAMP-dependent	402,0339	572	841.1746	632.0978	8.01E-01	2.45E-01	2 Unchanged Medium
P09131	F10gh1	Protein P3		629	901,5293	724.5168		-3.19E-02	2 Unchanged Medium
O60927	P02gh1	protein phosphatase 1; regulator	529.0069	607 1760	617.0621	584,2407	2.22E-01	1.986-01	2 Unchanged Medium
P41236	C08cd1	protein phosphatase 1; regulator		621	1404.955 583,7663		-2.99E-01	2.42E-02	2 Unchanged Medium
Q15435	D15ab8	protein phosphatase 1; regulator		963	753,9046	578,9436 784.0242	1.34E-01	2.23E-01	2 Unchanged Medium
P05323	E13ef6	protein phosphatase 2 (formerly:		1120	796,6526		2.48E-01 -4.10E-01	6.01E-01	2 Unchanged Medium
Q15257	G01ef5	protein phosphatase 2A; regulato		905	1160.437	996,8116		8.39E-02	2 Unchanged Medium
P33172	F01ab8	protein phosphatase 4 (formerly)		1220	1064,665		3.26E-01 -7.51E-01	-3.29E-02	2 Unchanged Medium
O00743	F03ab8	protein phosphatase 6; catalytic:	998.7	1070	738,4815		-4.35E-01	1.02E-01	2 Unchanged Medium
P38391	E21cd7	protein translocation complex bet		1050	1428,451	1175.447	4.45E-01		2 Unchanged Medium
P78324	M17cd4	protein tyrosine phosphatase; no		741	591.4877		-1.12E-01	2.13E-01	2 Unchanged Medium
Q13332	P21ab8	protein tyrosine phosphatase; rer		1880	2019.865	1928.827	1.02E-01	1.57E-03	2 Unchanged Medium 2 Unchanged Medium
Q9HAD6	C08gh6	protocadherin 16 dachsous-like (906	1070.494	981.1143		-9.50E-02	2 Unchanged Medium
O43445	E11cd5	PRP4 pre-mRNA processing fact	777,1081	549	569,3749		-4.49E-01		2 Unchanged Medium
Q9Y6B3	L19ef2	PTD013 protein	652.1251	701	1031.739	794.9108	6.62E-01	1.04E-01	2 Unchanged Medium
Q9HAN1	P19gh6	pumilio homolog 1 (Drosophila)	842.4925	1170	1047.18	1019.233	3.14E-01	4.71E-01	2 Unchanged Medium
Q9NQA4	C23ef4	pulative acid-sensing ion channe	742.723	762	678.3813	727.7123		3.70E-02	2 Unchanged Medium
043598	J05cd6	pulative c-Myc-responsive	473,6498	521	459.383		-4.41E-02	3.90E-01	2 Unchanged Medium
O43257	M12cd6	putative cyclin G1 interacting pro		1190	1238,925	1183.655	1.48E-01	9.40E-02	2 Unchanged Medium
Q9NY06	M07ef4	pulative integral membrane trans	966,4384	544	569,8565		-7.62E-01		2 Unchanged Medium
P17152	H21cd4	putative receptor protein	399.9305	650	591.3162	547.029	5.64E-01	7.00E-01	2 Unchanged Medium
Q9Y6I9	B13ef2	putative secreted protein	373.307	485	742,537	533.7168	9.92E-01	3.79E-01	2 Unchanged Medium
P54888	B12ab8	pyrroline-5-carboxylate syntheta:		302	905.8993	538,0842	1.16E+00	-4.31E-01	2 Unchanged Medium
Q13876	B03ef7	quiescin Q6	745.299	585	824.2489	718,0657	1.45E-01	-3.50E-01	2 Unchanged Medium
014679	K24ef5	quinone oxidoreductase homolog		1160	651,2257		-1.26E-01		2 Unchanged Medium
P53611	N16ab8	Rab geranylgeranyltransferase; t		727	938.6179	807,7755	3.08E-01	-6.14E-02	2 Unchanged Medium
P47224	D06ab8	RAB interacting factor	645,5902	859	577,3956		-1.61E-01	4.11E-01	2 Unchanged Medium
Q15907	H04cd4	RAB11B; member RAS oncogen		1030	1224.758	1060.693		1.53E-01	2 Unchanged Medium
P51153	D04ab8		619.0867	935		895,2783	8.70E-01		2 Unchanged Medium
P35287	J24ef2	RAB14; member RAS oncogene	543.7425	584	586,1635		1.08E-01		2 Unchanged Medium
Q9NP72	P02ef8		459,4786	794	634.6616	629.3713	4.66E-01		2 Unchanged Medium
P11476 Q9H0U4	H22ab8	RAB1A; member RAS oncogene	1039,455	767	993.586	933.4801	-6.51E-02		2 Unchanged Medium
P08886	L08gh7 B22ab8	RAB1B; member RAS oncogene RAB2; member RAS oncogene fr	1402 702	691	696.5292		4.02E-01		2 Unchanged Medium
095716	P04cd4	RAB3D; member RAS oncogene is	1403,703	1900	2195.304	1858.384	5.65E-01	3.54E-01	2 Unchanged Medium
P35239	D02ab8	RABSB; member RAS oncogene	500,9015	638	556,2242	701.181	-7.08E-01	5.10E-01	2 Unchanged Medium
P20340	C07ef6	RAB6A; member RAS oncogene	503 4507	708	393.3892	500,5498	-6.05E-01	2.42E-01	2 Unchanged Medium
P43487	D16ab8	RAN binding protein 1		655	416.7552		-2.72E-01		2 Unchanged Medium
Q9UI26	B20ef1	Ran binding protein 11	1438.305	1180		1468.585	3.13E-01		2 Unchanged Medium
P47736	D18ab8	RAP1; GTPase activating proteir	1505.224		1273.677	1319.368	-2.41E-01	-3.52E-01	2 Unchanged Medium
Q15382	F17cd2	Ras homolog enriched in brain 2	1435 507	414	461,3322		-9.25E-01		2 Unchanged Medium
P52199	B18ab2	ras homolog gene family; membe	534 E1E0	2190 1520	1635.792	1754.23	1.88E-01	6,1UE-U1	2 Unchanged Medium
P35238	B20ab2	ras homolog gene family; membe	1022 646	872	1010 477	971 4640	-4.22E-01	1,505+00	2 Unchanged Medium
Q14644	J08ef7	RAS p21 protein activator (GTPa	615 046	623	627.059	622 1475	-4.48E-03	•2.29E•U1	2 Unchanged Medium
Q15404	A10cd8	Ras suppressor protein 1	1543.953		2317.155	1925 644	2.58E-02	1./4E-02	2 Unchanged Medium
Q13283	H14cd5	Ras-GTPase-activating protein S	846 7267	789			5.86E-01	3.11E-U1	2 Unchanged Medium
Q9NVQ9	M05gh1	Rec8p; a meiolic recombination s	1334 371	1200	1325.85	1286 027	-5.68E-01 -9.24E-03	4 ESE 04	2 Unchanged Medium
O60895	K17cd6	receptor (calcitonin) activity modi	572 8174	510	899 2606	663 6562	6.51E-01	1 435 04	2 Unchanged Medium
		,, 111041		0.5		0002	J.J 1E*U1	-1.400-01	2 Unchanged Medium

P35244	122ef6	replication protein A3 (14kD)	544.0212	502	924.5385	656,7974	7.65E-01	-1.16F-01	2 Unchanged Medium
Q99623	D03cd7	repressor of estrogen receptor at	1139.283	689	1233.151	1020,486		-7.25E-01	2 Unchanged Medium
Q9UMQ4		ret proto-oncogene (multiple end	769,2484	602	803.931	725,1999	6.36E-02	-3.53E-01	2 Unchanged Medium
Q9NQC3		relicuton 4	1459.586	586	927.9776	991.1835		-1.32E+00	2 Unchanged Medium
P47804	P03cd1	retinal G protein coupled recepto		497	627.4176	509.1489	6.38E-01	3.02E-01	2 Unchanged Medium
Q16576 O95357	P07ef6 D02cd4	relinoblastoma binding protein 7	1539,006	1410	798.8744		-9.46E-01		2 Unchanged Medium
Q15299	D20ef1	retinoic acid induced 3 retinoic acid receptor-beta assoc	678.1926	713 979	208.6352 1013.444		-1.71E+00	7.14E-02	2 Unchanged Medium
Q9UMY1		retinoic acid repressible protein	1060,232 610,5279	726	586.2413		-6.51E-02		2 Unchanged Medium
Q13017	M15ef6	Rho GTPase activating protein 5	560.443	807	514.6138		-5.86E-02 -1.23E-01	2.51E-01 5.27E-01	2 Unchanged Medium
043182	B24ab2	Rho GTPase activating protein 6	1257.146	806	895.7191		-4.89E-01		2 Unchanged Medium 2 Unchanged Medium
060274	F04gh6	Rho-specific guanine nucleotide	651.9484	577	366.4761		-8.31E-01		2 Unchanged Medium
P07998	P21cd1	ribonuclease; RNase A family; 1	1294.476	3040	1592.811	1976.991	2.99E-01		2 Unchanged Medium
P04843	B12cd1	ribopherin I	1493,814	1360	2248.214	1700,316	5.90E-01	-1.37E-01	2 Unchanged Medium
P04844	B14cd1	ribophorin II	1574.181	1170	1880.047	1540.461	2.55E-01	-4.32E-01	2 Unchanged Medium
P26373 P40429	D11cd1 H12ef1	ribosomal protein L13	2091.256	1200	1924.546		-1.20E-01		2 Unchanged Medium
P39019	J05cd1	ribosomal protein L13a ribosomal protein S19	1928.024 1794.644	2060	1091.46		-8.21E-01	9.89E-02	2 Unchanged Medium
P78317	B04cd1	ring finger protein 4	756.2395	1480 661	2253.771 668.8027	1841.785	3.29E-01 -1.77E-01		2 Unchanged Medium
Q99942	F04cd2	ring finger protein 5	1016.837	949	1126.187	1030,595	1.47E-01		2 Unchanged Medium
043148	A15cd4	RNA (guanine-7-) methyltransfer.		730	390.498		-5.52E-01	3.50E-01	2 Unchanged Medium 2 Unchanged Medium
002916	F06ab8	RNA binding motif protein 4	533.7121	628	430,5298	530,87	-3.10E-01	2.36E-01	2 Unchanged Medium
Q9Y580	J10gh6	RNA binding motif protein 7	368.8423	710	558,043	545.5838	5.97E-01	9.45E-01	2 Unchanged Medium
Q14621	H07cd7	RNA binding protein (autoanliger	499.5699	737	506,822	581.142	2.08E-02		2 Unchanged Medium
Q93062	K17cd7	RNA-binding protein gene with m		659	640.4514		-4.00E-01		2 Unchanged Medium
Q99497 Q99584	B15cd7	RNA-binding protein regulatory s	1914.86	1400	2563.974	1958.111	4.21E-01		2 Unchanged Medium
P23526	L17cd2 A05ef1	S100 calcium binding protein At:	492.6196	421	675.1457	529.713		-2.25E-01	2 Unchanged Medium
043865	H21ab2	S-adenosylhomocysteine hydrola S-adenosylhomocysteine hydrola		603	478,692		-2.02E-01	1.32E-01	2 Unchanged Medium
014828	D02cd5	secretory carrier membrane prote		526 460	466,4887		-1.65E+00		2 Unchanged Medium
075326	L05cd3	sema domain; Immunoglobulin de		768	481.9262 943.1132	835.2276	-4.57E-01		2 Unchanged Medium
043278	116cd2	serine protease Inhibitor; Kunitz I		1260	641.0775		2.46E-01 -1.01E+00		2 Unchanged Medium
000271	C05ef5	serine protease inhibitor; Kunitz I		1740	876.4128		-6.55E-01	3.36E-01	2 Unchanged Medium
Q9UEW8	N22ef5	serine threonine kinase 39 (STE		1030	841.1872	1032.185	-5.48E-01		2 Unchanged Medium 2 Unchanged Medium
Q13043	P15cd2	serine/threonine kinase 4	750.7132	838	756.1662	781.5546	1.04E-02		2 Unchanged Medium
Q9Y282	F01ef2	serologically defined breast canc	1279.414	1160	1907.544	1448.986	5.76E-01		2 Unchanged Medium
P35542	A02gh1	serum amyloid A4; constitutive	754.5041	1310	811.5701	959,3499	1.05E-01	7.98E-01	2 Unchanged Medium
075790	122cd6	seven Iransmembrane domain pr		778	799.0998	779.5745	6.97E-02	3.16E-02	2 Unchanged Medium
Q9P0V3	H17ef3	SH3-domain binding protein 4	1204.005	1710	1539,363	1483.624	3.54E-01	5.04E-01	2 Unchanged Medium
Q99519	C13ab8	sialidase 1 (lysosomal sialidase)		680	498.7449		-5.15E-01		2 Unchanged Medium
Q92185 Q9Y6A9	J20cd1	sialyltransferase BA (alpha-N-acı		478	462.7941		-4.28E-01		2 Unchanged Medium
P21378	C05ef8 I04ef3	signal peptidase 12kDa signal peptidase complex (18kD)	449.125	742	887.4615	692.8575	9.83E-01	7.24E-01	2 Unchanged Medium
015302	P10ab3	signal recognition particle 72kD	1040.23	515 818	732.1592 841.5078	626.0198		-2.91E-01	2 Unchanged Medium
076094	J02cd2	signal recognition particle 72kD	1473.56	965	1422.2	899.9534 1286,822	-3.06E-01 -5.12E-02		2 Unchanged Medium
Q9UNL2	J10cd2	signal sequence receptor; gamm		450	1654.744	1085,128		-1.35E+00	2 Unchanged Medium 2 Unchanged Medium
P40763	K09ef5	signal transducer and activator o		545	348.0347		-9.48E-01		2 Unchanged Medium
Q92783	M24cd3	signal transducing adaptor molec	827.9851	1020	725.7017		-1.90E-01	3.01E-01	2 Unchanged Medium
Q9H723	N01gh5	similar to rat nuclear ubiquitous c	749,7661	530	1258.65	846.2392		-5.00E-01	2 Unchanged Medium
015258	A04cd7	similar to S. cerevisiae RER1	710.8499	635	933.6382	759.8427	3.93E-01	-1.63E-01	2 Unchanged Medium
O14834 O43617	A11cd7	similar to S. pombe dim1+	701.2292	753	903,8926	786,0877	3.66E-01	1.03E-01	2 Unchanged Medium
Q04837	H05ef3 G11cd2	similar to yeast BET3 (S. cerevis		1750	1443.578	1650,871	-2.85E-01		2 Unchanged Medium
P05455	E19gh1	single-stranded DNA binding pro Sjogren syndrome antigen B (aut		894	1034,75	1064.292	-2.89E-01	-5.00E-01	2 Unchanged Medium
P43331	K08cd2	small nuclear ribonucleoprotein (744 1260	930.6428 1042.335	841.9268 1073.82	1.29E-01		2 Unchanged Medium
P09012	M12cd2	small nuclear ribonucleoprotein p		310	685.6636	514.5708	1.86E-01 3.23E-01	4.63E-01	2 Unchanged Medium
P08579	A03cd2	small nuclear ribonucleoprotein ;	502.6355	570	669.9879	581.0279	4.15E-01	-8.22E-01 1.83E-01	2 Unchanged Medium
P09234	A05cd2	small nuclear ribonucleoprotein ;	1054.327	1450	1302.742	1269.64	3.05E-01	4.62E-01	2 Unchanged Medium 2 Unchanged Medium
P08578	A07cd2	small nuclear ribonucleoprotein p	1901,214	1740	1818.557	1819.876		-1.28E-01	2 Unchanged Medium
Q15357	A11cd2	small nuclear ribonucleoprotein p	382.4892	734	593.847	570.2015	6,35E-01	9.41E-01	2 Unchanged Medium
P14678	C23cd3	small nuclear ribonucleoprotein p	2036.574	1280	1737.398	1686.175	-2.29E-01		2 Unchanged Medium
Q13487	P08cd1 E17cd3	small nuclear RNA activaling cor		684	648.0121	616,6789	3.24E-01	4.03E-01	2 Unchanged Medium
P35326 P55854	H04cd2	small proline-rich protein 2A	290.6174	1030	839.3072	721.2318	1.53E+00		2 Unchanged Medium
P14648	P19cd5	SMT3 suppressor of mif two 3 ho SNRPN upstream reading frame	191./43/	658	962.4453	804.107		-2.67E-01	2 Unchanged Medium
P55011	L17cd1	solute carrier family 12 (sodium/s		2270 740	1670.964 517.2382	1839.36	7.90E-02 -1.94E-01		2 Unchanged Medium
O95258	P15cd4	solule carrier family 25 (mitochor	476 3075	518			2.79E-01	3.23E-01	2 Unchanged Medium
015431	A21ab5	solute carrier family 31 (copper b		1010	1769.026	1336.855		1.20E-01 -2.77E-01	2 Unchanged Medium
P04920	L04cd1	solute carrier family 4; anion excl		1560	1704.134	1624.765		-5.33E-02	2 Unchanged Medium 2 Unchanged Medium
Q01650	J11ef5	solute carrier family 7 (cationic as	832.6996	632	1032.106	832,1088		-3.99E-01	2 Unchanged Medium
Q07890	H23cd2	son of sevenless homolog 2 (Dro		568	680,8529	575.1528	5.14E-01	2.51E-01	2 Unchanged Medium
O60749	A19cd2	sorting nextn 2	364,9343	931	769.5782	688.5915	1.08E+00		2 Unchanged Medium
O60493	A09cd4	sorting nexin 3	1008.916	1350	1180.469		2.27E-01	4.22E-01	2 Unchanged Medium
Q9Y5X3	G06ef4	sorting nextn 5	676.0975	647	688,5568	670.591	2.63E-02	-6.32E-02	2 Unchanged Medium
Q01082	E09cd2	spectrin; beta; non-erythrocytic 1		679		545.8922			2 Unchanged Medium
P52788 P17947	M10cd2 C19cd2	spermine synthase	895,1994	854	770.002	839,7693	-2.17E-01	-6.78E-02	2 Unchanged Medium
Q14818	D10ef6	spleen focus forming virus (SFF)		1150	1646.278	1308.573	3.32E-01		2 Unchanged Medium
Q15427	K11cd6	splicing factor 1 splicing factor 3b; subunit 4; 49kt	547.0998	499 612	782.2929 540.0723	609.3662 530.9445		-1.34E-01	2 Unchanged Medium
Q15815	M06cd2	splicing factor; arginine/serine-ric		553	551.9082		2.92E-01	4.71E-01	2 Unchanged Medium
Q01130	H22cd1	splicing factor; arginine/serine-ric	553,5808	527	488.5465	522.9400	-4.20E-01 -1.80E-01	-9.10E-01	2 Unchanged Medium
P23152	H24cd1	splicing factor, arginine/serine-ric.	1547.474	1460	1428.273	1480.13	-1.16E-01	7.10E-02	2 Unchanged Medium 2 Unchanged Medium
Q13503	L22cd4	SRB7 suppressor of RNA polyme	712.8859	574	725.0591	670,5377	2.44E-02	-3.135-01	2 Unchanged Medium 2 Unchanged Medium
Q9P2R9	C23ef2	SRp25 nuclear protein	495,796	558	515.095	522.8318	5.51E-02	1.69E-01	2 Unchanged Medium
Q9H6I2	F07gh5	SRY (sex determining region Y)-I	774,5162	970	714.8666	819.674	-1.16E-01	3.24E-01	2 Unchanged Medium
Q9BT81	113gh8	SRY (sex determining region Y)-I	497.0608	530	550.1706	525.7297	1.46E-01		2 Unchanged Medium
									-

060526	M03cd6	STIP1 homology and U-Box cont.	706,6078	538	543.7559	596,1019	-3.78E-01	-3 93F-01	2 Unchanged Medium
Q9Y6X1	A21ef4	stress-associated endoplasmic re		1220	2201,647	1700.638	3.87E-01		2 Unchanged Medium
P31948	M15ef3	stress-Induced-phosphoprotein 1		549	517.5834	500.9139	2.47E-01		
P21912	E09gh1	succinate dehydrogenase compli		902	751.7357	785,3111	9.93E-02		2 Unchanged Medium
014521	H04cd1	succinate dehydrogenase compli		505					2 Unchanged Medium
					672.375	601.9355		-3.18E-01	2 Unchanged Medium
P53597	109gh1	succinate-CoA ligase; GDP-formi		491	553,293	582,7447			2 Unchanged Medium
095605	P15cd5	SUMO-1 activating enzyme subu		499	433.2742	569,4163	-8.42E-01		2 Unchanged Medium
Q16550	K01cd2	suppressor of Ty 4 homolog 1 (S		1440		1081.744	2.85E-01	8.24E-01	2 Unchanged Medium
Q15526	K07cd2	surfeit 1	468.3614	522	583,8097	524.7026	3.18E-01	1.56E-01	2 Unchanged Medium
043539	P02cd1	SWI/SNF related; matrix associa	1047.14	1280	1212.614	1178.519	2.12E-01	2.85E-01	2 Unchanged Medium
Q9UPX1	D23cd7	synaptopodin	683,4973	536	421.1841	546.8885	-6.98E-01	-3.51E-01	2 Unchanged Medium
095721	009cd5	synaptosomal-associated protein	830,7721	1030	686.6755	849.1506	-2.75E-01	3,10E-01	2 Unchanged Medium
043391	H09cd2	syndecan binding protein (synter	625.6962	640	379.1899	548.1569	-7.23E-01	3.17E-02	2 Unchanged Medium
Q15833	J04cd2	syntaxin binding protein 2	661,0175	556	840.6951	686.0183		-2.49E-01	2 Unchanged Medium
Q92804	O20cd3	TAF15 RNA polymerase II; TATA	873.5584	605	1063,778	847,4936	2.84E-01	-5.30E-01	2 Unchanged Medium
Q9Y490	B04cd2	talin 1	1342,238	1170	2365,625	1625.187		-2.01E-01	2 Unchanged Medium
094797	H20ef7	Tara-like proteln	899,0807	448	496.1404	614.3777			2 Unchanged Medium
Q13311	C05cd6	Tax1 (human T-cell leukemia vin		1190	926.2314	1004.505	4.15E-02	4.00E-01	2 Unchanged Medium
Q16650	F08cd6	T-box; brain; 1	1407.309	1750	1777.889	1645.184		3.15E-01	
P51864	O04ef7	teratocardnoma-derived growth i		1590			3.37E-01		2 Unchanged Medium
P55061					1447.46	1690.282			2 Unchanged Medium
	019cd2	testis enhanced gene transcript (1330	1972,323	1615.143		-2.10E-01	2 Unchanged Medium
Q15569	P21cd2	lestis-specific kinase 1	837.6269	796	708.1178		-2.42E-01		2 Unchanged Medium
060636	F08cd5	tetraspan 2	468,1664	702	375.3225	515.2157	-3.19E-01	5.85E-01	2 Unchanged Medium
060637	F06cd5	tetraspan 3	1025,362	742			-9.09E-01		2 Unchanged Medium
095857	F13ef3	tetraspan NET-6 protein	702,1579	676	359.1921	578.9636	-9.67E-01	-5.58E-02	2 Unchanged Medium
Q99614	M03cd3	letratricopeptide repeat domain 1		891	635.5379	744.2616	-1.52E-01	3.35E-01	2 Unchanged Medium
Q1311B	LG5cd2	TGFB inducible early growth resp	445.9691	601	679.2931	575.4732	6.07E-01	4.31E-01	2 Unchanged Medium
Q15583	A16cd2	TGFB-induced factor (TALE fami	885.9	1290	1140,439	1105.268	3.64E-01	5,42E-01	2 Unchanged Medium
Q9H3N1	N03gh7	thioredoxin domain-containing	409.6333	426	686.7997	507,428	7.46E-01	5.60E-02	2 Unchanged Medium
043396	017cd5	thioredoxin-like; 32kD	1821,595	1710	1548,596	1693.179		-9.18E-02	2 Unchanged Medium
P04216	B02cd2	Thy-1 cell surface anligen	169.8064	298	4157,367	1541.609		8,10E-01	
P12956	K10ef6	thyroid autoantigen 70kD (Ku ani	1172.56	1490					2 Unchanged Medium
P16035	P11ef7				1395.803	1353.001	2.51E-01	3.46E-01	2 Unchanged Medium
		tissue inhibitor of metalloproteina		354	913.1351	542.4517		-2.54E-02	2 Unchanged Medium
Q9H2X8	108gh8	TLH29 protein precursor	710.2447	1000	1386,102	1033.648	9.65E-01	5.00E-01	2 Unchanged Medium
Q13077	L11cd2	TNF receptor-associated factor 1		1270	756.401	1097.404	-7.42E-01	6.49E-03	2 Unchanged Medium
Q9NZ34	P08gh4	TPA regulated locus	1074.319	1090	1284.44	1148.575	2.58E-01	1.69E-02	2 Unchanged Medium
Q15369	D18ef6	transcription elongation factor B :	771.5598	1100	693.1767	854.8396	-1.55E-01	5.11E-01	2 Unchanged Medium
Q15906	P11ef6	transcription factor-like 1	436.5561	510	878,9083	608.5936	1.01E+00	2.25E-01	2 Unchanged Medium
P50616	H08cd5	transducer of ERBB2; 1	888,8059	874	518.1816	760.3803	-7.78E-01	-2.40F-02	2 Unchanged Medium
Q13595	N04ef7	transformer-2 alpha (hira-2 alpha	247.4437	612	698,379		1.50E+00		2 Unchanged Medium
P37802	F05cd3	transgelin 2	1986,716	2240	1173,434		-7.60E-01	1.70E-01	2 Unchanged Medium
Q9Y5Z9	D01cd8	transitional epithelia response pr		875	1126,793		-2.33E-01		
Q99595	K10cd6	transfocase of inner mitochondria		665	448.5161				2 Unchanged Medium
060830	A09cd6	translocase of inner mitochondria					-6.87E-03	5.60E-01	2 Unchanged Medium
014925				1590	1992.925	1744.175		-4.77E-02	2 Unchanged Medium
	124cd6	translocase of inner mitochondria		813	653,4301		-1.91E-01	1.25E-01	2 Unchanged Medium
Q9Y5J9	E08cd8	translocase of inner milochondria		1550	1653.524	1615.991		-7.98E-02	2 Unchanged Medium
Q15629	G18ef3	translocating chain-associating n		1060	624.4499	897.0752	-6.84E-01	8.48E-02	2 Unchanged Medium
Q99442	C10cd2	translocation protein 1	526.8783	772	1065.397	788.001	1.02E+00	5.51E-01	2 Unchanged Medium
Q99805	A18cd5	transmembrane 9 superfamily me		1260	957.2801	1081.028	-9.63E-02	3.03E-01	2 Unchanged Medium
Q9Y2B0	C20ef3	transmembrane protein 4	681.3857	1320	985.8397	995.6424	5,33E-01	9.54E-01	2 Unchanged Medium
P49755	J05ef4	transmembrane trafficking proteir	1311.203	469	700.7243	827.066	-9.04E-01	-1.48E+00	2 Unchanged Medium
	F03gh4	transmembrane; prostate androg	1287.98	841	662,4165		-9.59E-01		2 Unchanged Medium
Q03519	B07cd1	transporter 2; ATP-binding casse	851,2148	573	680.8519	701.534	-3.22E-01		2 Unchanged Medium
Q9NWF9	P02gh2	TRIAD3 protein	665,9772	444	688.6646	599.4053		-5.86E-01	2 Unchanged Medium
Q15656	007ef7	TRK-fused gene .	1878,254	1640	1858.307		-1.54E-02		2 Unchanged Medium
Q92734	J17cd7	TRK-fused gene	1450,307	1810	1609,067	1624.429	1.50E-01	3.23E-01	2 Unchanged Medium
Q12815	A21ef7	trophinin associated protein (tast			722.8644				
P09493	002cd1	tropomyosin 1 (alpha)		429			-1.85E-01		
P49411			988,1694		1004.982	807.373	2.43E-02		2 Unchanged Medium
P05217	M11cd3 F03gh6	Tu translation elongation factor; i tubulin; beta; 2		555	476.4592 920.1831			7.90E-02	2 Unchanged Medium
			1203.686	705			-3.87E-01		2 Unchanged Medium
P23258	K03cd3	tubulin; gamma 1	1733.971	1220	2149.365	1702.331	3,10E-01		2 Unchanged Medium
Q9UBN6	A16el6	tumor necrosis factor receptor su		751	531.524	584.4772	1.75E-01	6.74E-01	2 Unchanged Medium
Q13829	L22ef4	tumor necrosis factor; alpha-indu	849,276	792	552.7776		-6.20E-01		2 Unchanged Medium
000496	K21cd3	turnor suppressing subtransferab		554	800.8751		3.50E-01		2 Unchanged Medium
Q99816	G04cd4	turnor susceptibility gene 101	723.8133	766	627.1295	705.4904	-2.07E-01	8.08E-02	2 Unchanged Medium
P16422	C03ab7	tumor-associated calcium signal	1448.512	791	373,663		-1.95E+00		2 Unchanged Medium
Q9NP84	K03ef2	type I transmembrane protein Fn	691.89	498	560.2824		-3.04E-01		2 Unchanged Medium
Q9UJ47	009cd8	type i transmembrane receptor (s	792,9873	819	809.9056	807.3703	3.05E-02		2 Unchanged Medium
P31946	K21ef6	tyrosine 3-monooxygenase/tryptc			724.9898		-7.52E-01		2 Unchanged Medium
P35214	J19ef4	tyrosine 3-monooxygenase/trypto		989	682.5327		-3.68E-01		2 Unchanged Medium
P27348	G15cd7	tyrosine 3-monooxygenase/tryptc		1540	1104.855		-9.91E-01		
060704	J01cd3	tyrosylprotein sulfotransferase 2		299					2 Unchanged Medium
Q9Y4Z0	G09cd8					547.7036			2 Unchanged Medium
		U6 snRNA-associated Sm-like pr		689	914.9547				2 Unchanged Medium
P14927	G06cd4	ubiquinol-cytochrome c reductas		817			-6.04E-02		2 Unchanged Medium
Q9U102	PtBef7	ubiquinol-cytochrome c reductas			1546,568		6.97E-01		2 Unchanged Medium
P31930	A22cd3	ubiquinol-cytochrome c reductas		849	686.9864	794.0747	-3.01E-01	3.33E-03	2 Unchanged Medium
P22695	A24cd3	ubiquinol-cytochrome c reductas:	468,5936	636	581,791	562.0633		4.40E-01	2 Unchanged Medium
P07919	E08cd4	ubiquinol-cytochrome c reductas-		1810	1005.378		-4.02E-01	4,49E-01	2 Unchanged Medium
P47985	E06cd4	ublquinol-cytochrome c reductas				1306.086			2 Unchanged Medium
Q9NZ09	N17ef1	ubiquitin associated protein	1588.177			1598.619			2 Unchanged Medium
Q9UNP0		ubiquitin specific protease 15	525.3364	492	597,7817	538.3523	1.865-01	-9 ARE-02	2 Unchanged Medium
Q9UHP3		ubiquitin specific protease 25	554.0657	724	561.5817		1.94E-02		2 Unchanged Medium
P22314	O07cd3	ubiquitin-activating enzyme E1 (/		1640		1788.503			
Q13404	O23cd3	ubiquitin-conjugating enzyme E2			1129.000	4420 555	4.71E-03	-1./3E-UI	2 Unchanged Medium
P51669	D05cd4		1266.25	1300		1138.559			2 Unchanged Medium
P47986	011cd3	ubiquitin-conjugating enzyme E2			1173.73	1247.427	-1.09E-01	4.05E-02	2 Unchanged Medium
1 41 300	0.100	ubiquitin-conjugating enzyme E2	010,3009	320	307.7317	567.8765	-1.19E-01	-2.47E-01	2 Unchanged Medium

O14933	H14cd4	ublquitin-conjugating enzyme E2	717,4244	893	596,6415	735.6835	-2.66E-01	3,16E-01	2 Unchanged Medium
Q16781	D07cd4	ubiquitin-conjugating enzyme E2		816	533,7389			-1.78E-01	2 Unchanged Medium
Q93068	A02cd3	ublquitin-like 1 (sentrin)	1376,087	1530	1294,307		-8.84E-02		2 Unchanged Medium
Q15386	l10gh1	ubiquitin-protein isopeptide ligas	531.8659	630	628.2454	596.8686		2.45E-01	2 Unchanged Medium
Q9UBK9	115cd4	ubiquitously-expressed transcript		1060	1384,764	1125,743	5.65E-01	1.74E-01	2 Unchanged Medium
060512	Ettab3	UDP-Gal:betaGlcNAc beta 1:4- c	436,7634	610	542.39	529.5839	3.12E-01	4.81E-01	2 Unchanged Medium
P78383	P20cd5	UDP-galactose transporter relate		1700	2180,117	1816,963	4.74E-01	1.15E-01	2 Unchanged Medium
Q16851	124cd4	UDP-glucose pyrophosphorylase	1324.08	1260	1318,78	1302,468		-6.64E-02	
Q9NZ45	L02gh3	uncharacterized hematopoletic st	652.0685	670	696,5692	672,9895	9.52E-02	3.98E-02	2 Unchanged Medium
Q9NZ32	L20gh4	uncharacterized hypothalamus p	1035,575	1250	1354,231	1214,412	3.87E-01	2.75E-01	2 Unchanged Medium
Q9NZ29	L16gh4	uncharacterized hypothalamus p	659.53	992	1118,776	923,518	7.62E-01	5.89E-01	2 Unchanged Medium
Q15853	C02cd3	upstream transcription factor 2; c	601.6839	635	698.0128	644.9023	2.14E-01	7.78E-02	2 Unchanged Medium
Q92528	D02ef7	uridine monophosphate kinase	560,3976	685	623.5243	623.0076	1.54E-01	2.90E-01	2 Unchanged Medium
P15692	M04ef7	vascular endothelial growth facto	460.9257	539	857,3853	619,202	8.95E-01	2.27E-01	2 Unchanged Medium
P50552	C08cd3	vasodilator-stimulated phosphopi	468.4573	648	392,5569	502.9122	-2.55E-01	4.67E-01	2 Unchanged Medium
060763	H16cd3	vesicle docking protein p115	639.538	543	745.4806	642.7624	2.21E-01	-2.35E-01	2 Unchanged Medium
Q15836	A15ef5	vesicle-associated membrane pri	804,2648	1370	1060.211	1076,578	3.99E-01	7.63E-01	2 Unchanged Medium
Q9UEU0	022cd6	vesicle-associated soluble NSF ε	468,6024	697	698,3534	621,1953	5.76E-01	5.72F-01	2 Unchanged Medium
P14921	E24ef5	v-els erythroblastosis virus E26 c	913.893	765	291,1538	656,6528	-1.65E+00	-2.57E-01	2 Unchanged Medium
P18206	C14cd3	vinculin	580.3315	857	865,7505		5.77E-01	5.62E-01	2 Unchanged Medium
P00540	G08ef5	v-mos Moloney murine sarcoma .	1115,145	728	663,6747			-6.16E-01	2 Unchanged Medium
P01106	E18ef5	v-myc myelocytomatosis viral on:	537.8131	601	405.1047	514.7639		1.61E-01	2 Unchanged Medium
P21796	C16cd3	voltage-dependent anion channe		510	584,201	525,9036		7.47E-02	2 Unchanged Medium
Q15765	C12cd3	von Hippel-Lindau binding protei	689,743	705	891.2621	762,0788	3.70E-01	3.20E-02	2 Unchanged Medium
P10398	D01ab4	v-raf murine sarcoma 3611 viral c	1061.812	639	532.2153	744,4583	-9.96E-01	-7.32E-01	2 Unchanged Medium
P11233	A19ef6	v-ral simian leukemia viral oncog	848.7099	1060	1081,428	995.8044	3.50E-01	3.17E-01	2 Unchanged Medium
Q9Y6W5	A22gh1	WAS protein family; member 2	1283.077	1630	1493,145	1468.514	2.19E-01	3,45E-01	2 Unchanged Medium
Q9UPY6	L06cd6	WAS protein family; member 3	1503.584	1210	2151.643	1623,108	5.17E-01	-3.09E-01	2 Unchanged Medium
075083	D15cd5	WD repeat domain 1	892.4425	977	963.3448	944,1492	1.10E-01	1.30E-01	2 Unchanged Medium
076071	A24cd5	WD40 protein Clao1	609.8821	898	730.5311	746,1814	2.60E-01	5.58E-01	2 Unchanged Medium
Q9HCN4	H11gh6	XPA binding protein 1; putative A	1199.21	1070	507.3277	923.9753	-1,24E+00	-1.71E-01	2 Unchanged Medium
Q9UM05	N17ef3	yeast Sec31p homolog	.1887.447	1590	1542.778	1672.59	-2.91E-01	-2.50E-01	2 Unchanged Medium
Q9UQR1	P06gh1	zinc finger protein 148 (pHZ-52)	513.8639	737	563,7625	604.9201	1.34E-01	5.21E-01	2 Unchanged Medium
Q14119	G21ef5	zinc finger protein 161	1199.558	1760	1197.34	1387.253	-2.67E-03	5.57E-01	2 Unchanged Medium
Q9NW07	F06gh2	zinc finger protein 358	402.8342	575	684,7069	554.3431	7.65E-01	5.15E-01	2 Unchanged Medium
P20694	D04ef6	zinc finger prolein 9 (a cellular re	1042.65	1010	856,5924	969,9823	-2.84E-01	-4.49E-02	2 Unchanged Medium

WHAT IS CLAIMED IS:

1. A method for transforming IP cells that express markers of acinar cells and liver-associated genes into insulin-producing cells in vitro, comprising culturing said IP cells in a cell culture medium comprising an effective amount of at least one differentiation promoting factor selected from the group consisting of C-Natriuretic Peptide (CNP), Calcitonin Gene Related Peptide, Cholera Toxin B Subunit, Dexamethasone, Gastrin-Releasing Peptide, Laminin, Met-Enkephalin, PDGFAA+PDGFBB, Sonic Hedgehog, and Substance P such that the IP cells are transformed into insulin-producing cells.

- 2. The method of claim 1, wherein the IP cells are derived from a culture of pancreatic acinar cells.
- 3. The method of claim 2, wherein the cells are human.
- 4. The method of claim 1, further comprising contacting said cells with a substrate that is coated with one or more extracellular matrix molecules.
- 5. The method of claim 4, wherein the extracellular matrix molecules are collagen I, collagen VI, collagen IV, vitronectin, and/or fibronectin.
- 6. The method of claim 4, wherein the substrate is on the surface of a flask, petri dish, plate, well or roller bottle, or is part of a scaffold.
- 7. The method of claim 1, wherein the medium is serum-free.
- 8. The method of claim 1, wherein the medium comprises serum.
- 9. The method of claim 7, wherein the medium comprises BSA, insulin, transferrin, selenium and epidermal growth factor (EGF).
- 10. The method of claim 3, wherein the cells are seeded on the substrate at a density of 5×10^3 to 20×10^5 cells/cm².
- 11. An isolated insulin-producing cell generated by the method of claim 1.
- 12. An insulin-producing cell, prepared by differentiating a mammalian acinar cell in vitro, wherein said insulin-producing cell has an expression profile after 16 days ex vivo as shown in Table 6.
- 13. A serum-free medium comprising at least one differentiation promoting factor selected from the group consisting of C-Natriuretic Peptide (CNP), Calcitonin Gene Related Peptide, Cholera Toxin B Subunit, Dexamethasone, Gastrin-

Releasing Peptide, Laminin, Met-Enkephalin, PDGFAA+PDGFBB, Sonic Hedgehog, and Substance P wherein said medium facilitates differentiation of IP cells into insulin-producing cells.

- 14. A serum free medium comprising a 1:1 mixture of DMEM and Hams F12 plus the components listed in Table 2.
- 15. A kit suitable for differentiating IP cells to insulin-producing cells, comprising
 - a) a base medium suitable for the cultivation of mammalian epithelial cells;
 - b) a collagen I coated culture substrate, and, separately packaged,
 - c) a serum-free medium supplement containing BSA, C-Natriuretic Peptide (CNP), Calcitonin Gene Related Peptide, Cholera Toxin B Subunit, Dexamethasone, Gastrin-Releasing Peptide, Laminin, Met-Enkephalin, PDGFAA+PDGFBB, Sonic Hedgehog, and Substance P or two or more of these components in combination, in suitable amounts to yield final concentrations in the completed medium as indicated in Table 1 herein.
- 16. The kit of claim 15, wherein the cell culture substrate is contained on the surface of a flask, bottle, petri dish, plate or well suitable for cell culture.
- 17. The method of claim 1, wherein the cell culture medium comprises a 1:1 mixture of DMEM and Hams 12.
- 18. The serum-free medium of claim 13 which comprises a 1:1 mixture of DMEM and Hams 12.
- 19. The method of claim 1, wherein said differentiation promoting factors have the concentrations in the medium as indicated in Table 1.
- 20. The serum-free medium of claim 13, wherein said differentiation promoting factors have the concentrations in the medium as indicated in Table 1.

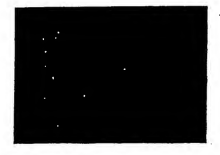


Figure 1A

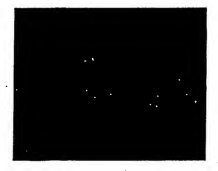


Figure 1B

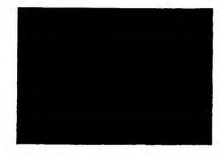


Figure 1C

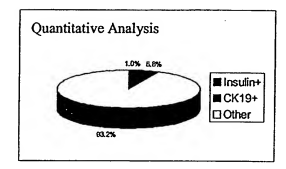
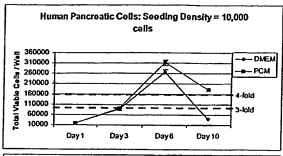
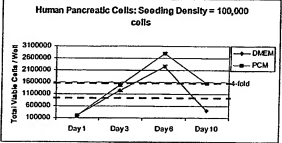


Figure 1D





Human pancreatic cells were seeded onto tissue culture treated polystyrene in either DMEM commercial media + 10% fetal bovine serum or in our BDT pancreatic cell media (PCM) + 10% fetal bovine serum. Replicate cultures were harvested at 3-day intervals via trypsinization and live cells (as determined by trypan blue exclusion) were enumerated on a hemacytometer.

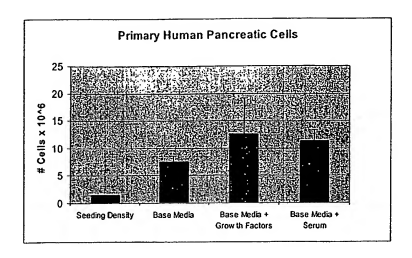


Figure 4A

Total Cell Number: Effect of Surface

Figure 4B

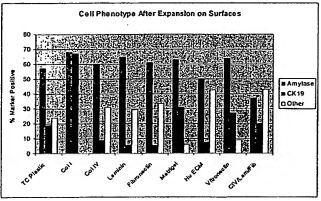


Figure 5A

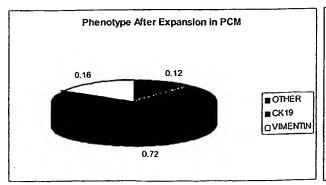
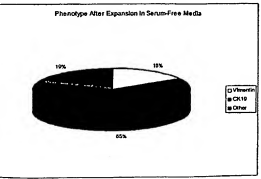
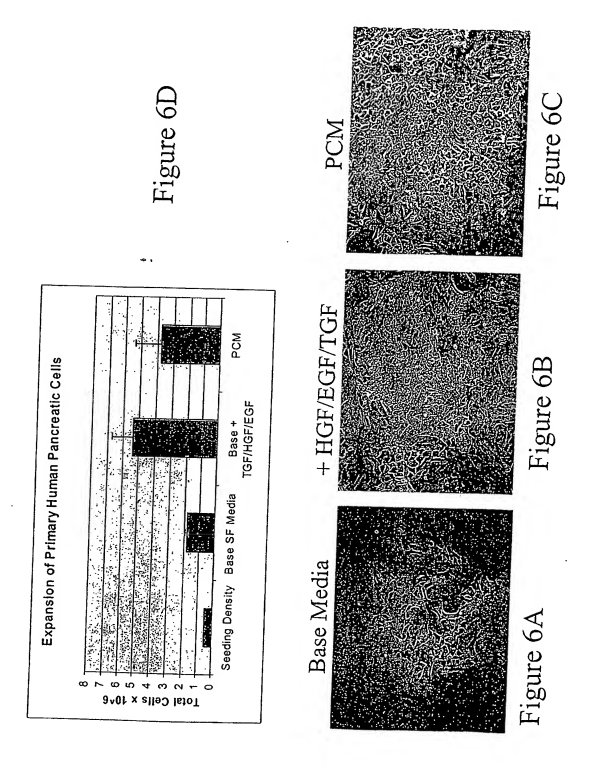


Figure 5B





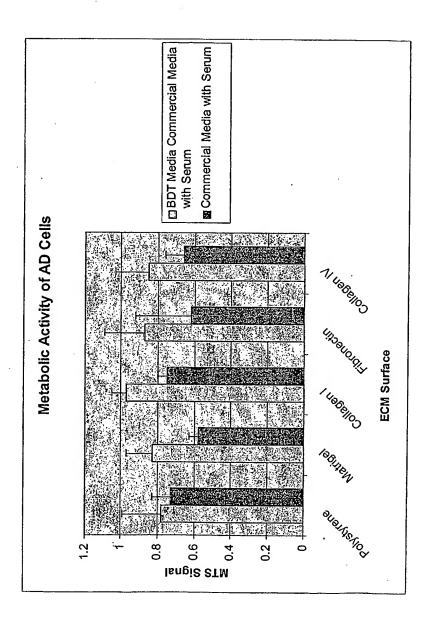


Figure 7

Figure 8A

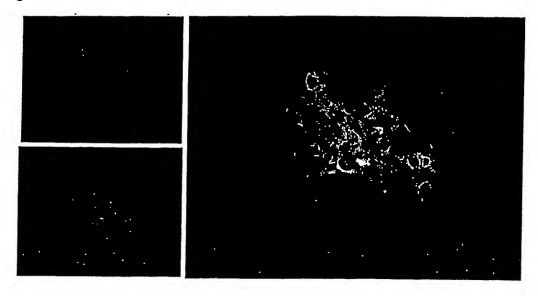


Figure 8B Figure 8C

Figure 9

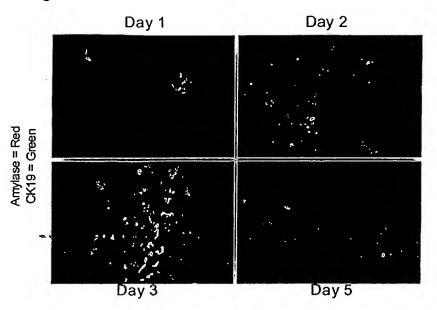
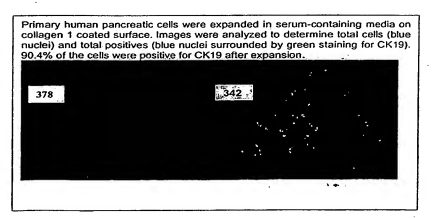


Figure 10A-B



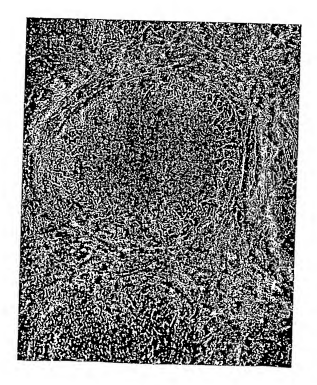
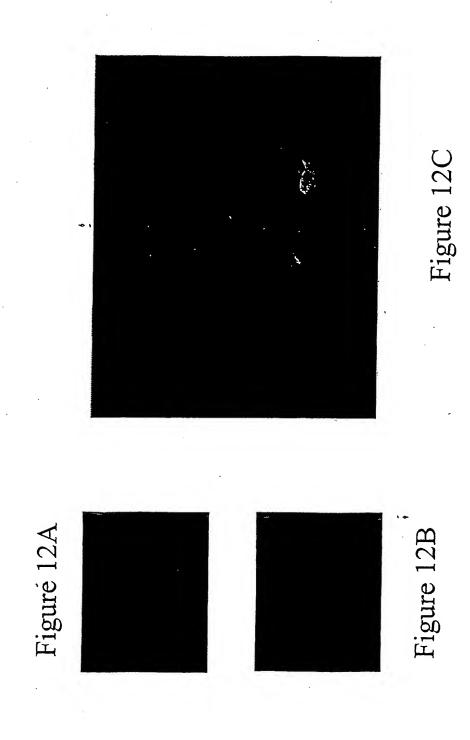
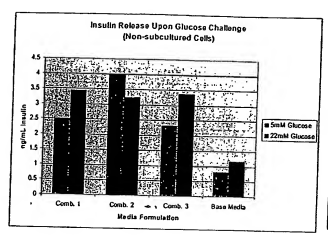
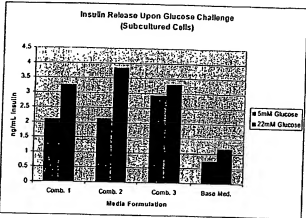


Figure 11







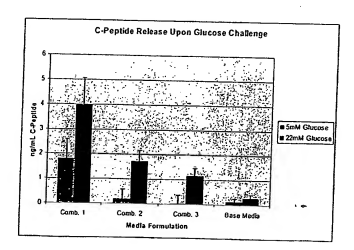
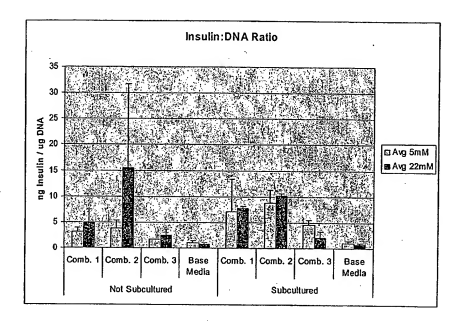


Figure 13C



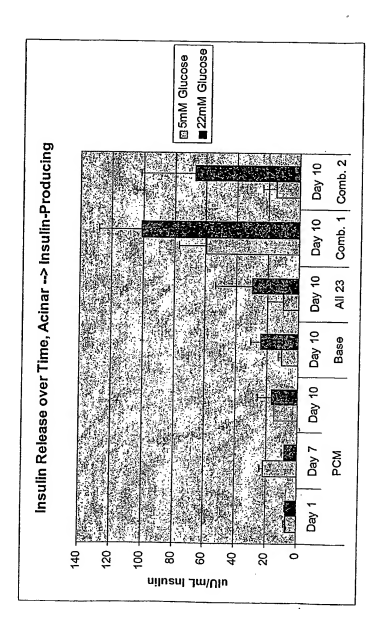
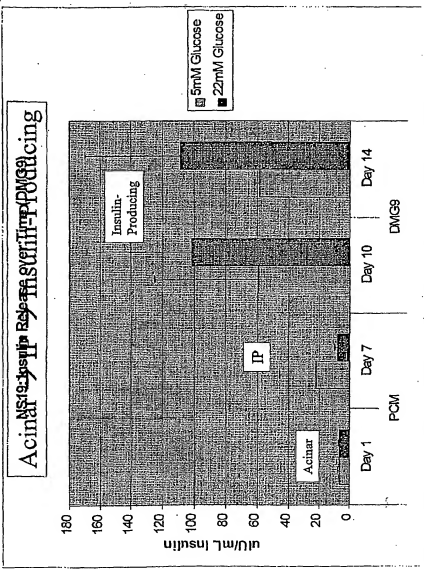
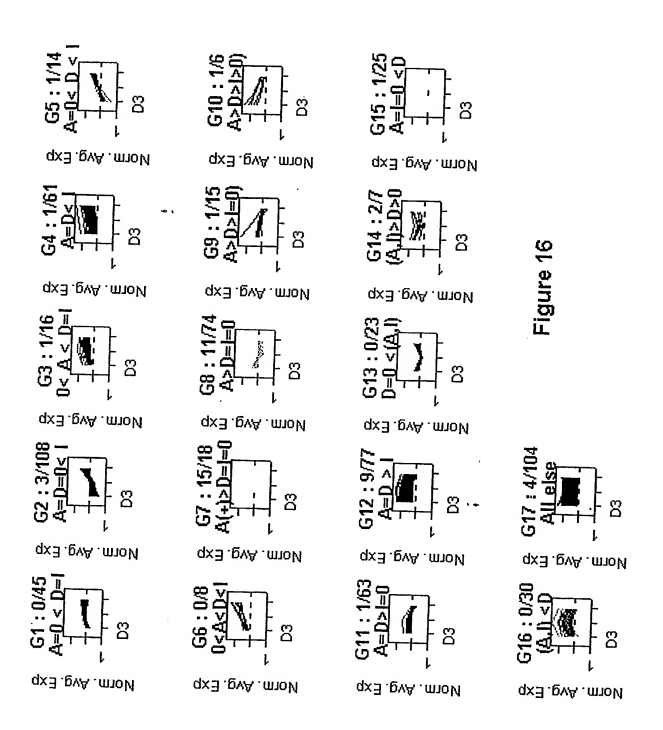


Figure 15



7, thus generating a culture of IP cells at Day 7. On Day 7, the IP cells were washed and the PCM media was replaced with the G09 differentiation media containing the 30 factors listed in Table 2. At each time point, insulin release was measured by washing the cultures 3x with PBS, then challenging the cultures with a 1:1 mixture of DMEM and HAMs F12 containing either 5mM or 22 mM glucose. After 18 hours of Figure 15: Human pancreatic acinar cells were cultured on collagen 1 surface in PCM from Day 1 to Day exposure to the glucose, supernatants were collected and insulin measured by ELISA.

Figure 15A



14/14

INTERNATIONAL SEARCH REPORT

International application No.

PCT/US03/16096

A. CLAS	SSIFICATION OF SUBJECT MATTER : C12N 5/00,5/02, 15/00							
US CL	: 435/1.2, 70.1,70.3,325,366, 375, 373, 374, 3							
According to	International Patent Classification (IPC) or to both	national classification and IPC						
B. FIEL	DS SEARCHED							
Minimum do U.S.: 4	Minimum documentation searched (classification system followed by classification symbols) U.S.: 435/1.2, 70.1,70.3,325,366, 375, 373, 374, 377, 378, 383, 384, 385,388395,404,405							
Documentation	on searched other than minimum documentation to th	e extent that such documents are included	in the fields searched					
	ata base consulted during the international search (national search (national)	me of data base and, where practicable, so	earch terms used)					
	UMENTS CONSIDERED TO BE RELEVANT							
Category *	Citation of document, with indication, where a	``	Relevant to claim No.					
A,P	US 6,436,704 B1 (ROBERTS et al) 20 August 2002	2 (20.08.2002), see entire documnet.	1-20					
A	US 4,332,893 A (ROSENBERG) 01 June 1982 (01	.06.1982), see entire document.	1-20					
A,E	US 2003/011330 A1 (PERFETTI) 19 June 2003 (0:	1.06.2003), see entire document.	1-20					
A	US 2002/0015696 A1 (GERMAN) 07 February 200	02 (07.02.2002), see entire document.	1-22					
A,P	US 2003/0087394 A1 (SHARMA) 08 May 2003 (08.05.2003), see entire document 1-20							
A,P	US 2002/0072115 A1 (HARRISON et al) 13 June 2002 (13.06.2002), see entire 1-20 document.							
Α	US 2003/0138951 A1 (YIN) 24 July 2003 (24.07.20	003), see entire document.	1-20					
Further	documents are listed in the continuation of Box C.	See patent family annex.						
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